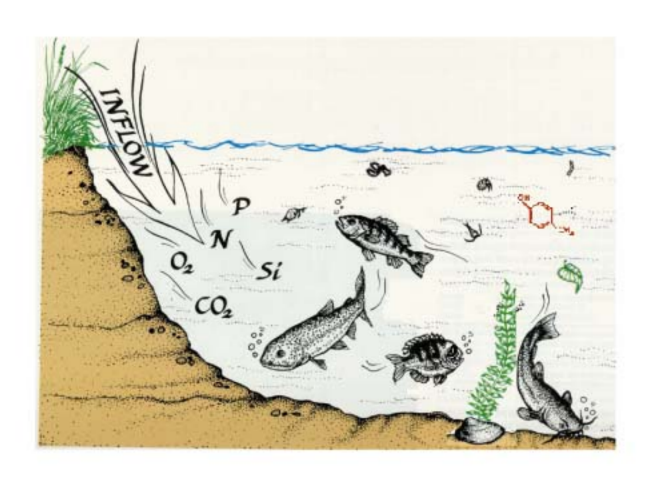


SEPA AQUATOX FOR WINDOWS

A MODULAR FATE AND EFFECTS MODEL FOR AQUATIC ECOSYSTEMS

RELEASE 1

VOLUME 2: TECHNICAL DOCUMENTATION



AQUATOX FOR WINDOWS

A MODULAR FATE AND EFFECTS MODEL FOR AQUATIC ECOSYSTEMS

RELEASE 1

VOLUME 2: TECHNICAL DOCUMENTATION

SEPTEMBER 2000

U.S. ENVIRONMENTAL PROTECTION AGENCY
OFFICE OF WATER
OFFICE OF SCIENCE AND TECHNOLOGY
WASHINGTON DC 20460

DISCLAIMER

This document has been approved for publication by the Office of Science and Technology, Office of Water, U.S. Environmental Protection Agency. Mention of trade names, commercial products or organizations does not imply endorsement or recommendation for use.

This document describes a new aquatic ecosystem simulation model. It is not intended to serve as guidance or regulation, nor is the use of this model in any way required. This document cannot impose legally binding requirements on EPA, States, Tribes, or the regulated community.

ACKNOWLEDGMENTS

This model has been developed and documented by Dr. Richard A. Park of Eco Modeling; most of the programming has been by Jonathan S. Clough under subcontract to Eco Modeling. It was funded with Federal funds from the U.S. Environmental Protection Agency, Office of Science and Technology under contract number 68-C4-0051 to The Cadmus Group, Inc.. Work assignment managers for The Cadmus Group have been Paul Jacobson, Jonathan Butcher, and William Warren-Hicks; their help in expediting the contractual arrangements and in reviewing the scientific approaches is appreciated. Revision of the documentation has been performed under subcontract to AQUA TERRA Consultants, Anthony Donigian, Work Assignment Manager, under EPA Contract 68-C-98-010.

Additional Federal funding for program development has come from the U.S. Environmental Protection Agency, Office of Pollution Prevention and Toxics, through Purchase Orders 7W-0227-NASA and 7W-4330-NALX to Eco Modeling.

The assistance, advice, and comments of the EPA work assignment manager, Marjorie Coombs Wellman of the Exposure Assessment Branch, Office of Science and Technology has been of great value in developing this model and preparing this report. Further technical and financial support from David A. Mauriello and Rufus Morison of the Office of Pollution Prevention and Toxics is gratefully acknowledged..

In an earlier version of the model developed at Abt Associates, Brad Firlie facilitated the programming; Rodolfo Camacho developed and programmed the inorganic sediment constructs; and review was provided by Lisa Akeson, Elizabeth Fechner-Levy, and Keith Sappington.

TABLE OF CONTENTS

1.1 Overview	1 1 3
SIMULATION MODELING 2 - 2.1 Temporal and Spatial Resolution and Numerical Stability 2 - 2.2 Uncertainty Analysis 2 - PHYSICAL CHARACTERISTICS 3 - 3.1 Morphometry 3 -	1 1 3
2.1 Temporal and Spatial Resolution and Numerical Stability2 -2.2 Uncertainty Analysis2 -PHYSICAL CHARACTERISTICS3 -3.1 Morphometry3 -	1 3
2.1 Temporal and Spatial Resolution and Numerical Stability2 -2.2 Uncertainty Analysis2 -PHYSICAL CHARACTERISTICS3 -3.1 Morphometry3 -	1 3
2.2 Uncertainty Analysis 2 - PHYSICAL CHARACTERISTICS 3 - 3.1 Morphometry 3 -	3
PHYSICAL CHARACTERISTICS 3 - 3.1 Morphometry 3 -	1
3.1 Morphometry	
3.1 Morphometry	
± •	
Volume	
Bathymetric Approximations	
3.6 Wind 3 -	14
RIOTA 4 -	1
e	
· · · · · · · · · · · · · · · · · · ·	
± •	
± •	
· · · · · · · · · · · · · · · · · · ·	
1 TOMOUOH	
REMINERALIZATION	1
	3.2 Washout 3 - 3.3 Stratification and Mixing 3 - 3.4 Temperature 3 - 3.5 Light 3 - 3.6 Wind 3 - BIOTA 4 - 4.1 Algae 4 - Light Limitation 4 - Nutrient Limitation 4 - Adjustment for Suboptimal Temperature 4 - Algal Respiration 4 - Photorespiration 4 - Algal Mortality 4 - Sinking 4 - Washout and Entrainment 4 - Chlorophyll a 4 - 4.2 Macrophytes 4 - 4.3 Animals 4 - Consumption, Defecation, and Predation 4 - Respiration 4 - Excretion 4 - Nonpredatory Mortality 4 - Gamete Loss and Recruitment 4 - Washout and Drift 4 - Vertical Migration 4 - Promotion 4 - REMINERALIZATION 5 - 5.1 Detritus 5 - Colonization 5 -

Decomposition
Sedimentation
5.2 Nitrogen 5 - 10
Assimilation
Nitrification and Denitrification 5 - 13
5.3 Phosphorus
5.4 Dissolved Oxygen
5.5 Inorganic Carbon
6. INORGANIC SEDIMENTS
6.1 Deposition and Scour of Silt and Clay
6.2 Scour, Deposition and Transport of Sand
6.3 Suspended Inorganic Sediments in Standing Water 6 - 7
7. TOXIC ORGANIC CHEMICALS
7.1 Ionization
7.2 Hydrolysis
7.3 Photolysis
7.4 Microbial Degradation
7.5 Volatilization
7.6 Partition Coefficients
7.7 Nonequilibrium Kinetics
Sorption and Desorption to Sedimented Detritus
Bioconcentration in Macrophytes and Algae
Macrophytes
Algae 7 - 27
Bioaccumulation in Animals
Gill Sorption
Dietary Uptake
Elimination
Linkages to Detrital Compartments
8. ECOTOXICOLOGY 8 - 1
8.1 Acute Toxicity of Compounds
8.2 Chronic Toxicity
9. REFERENCES
APPENDIX A. GLOSSARY OF TERMS A - 1
APPENDIX B. USER-SUPPLIED PARAMETERS AND DATA

PREFACE

The Clean Water Act—formally the Federal Water Pollution Control Act Amendments of 1972 (Public Law 92-50), and subsequent amendments in 1977, 1979, 1980, 1981, 1983, and 1987– calls for the identification, control, and prevention of pollution of the nation's waters. In the National Water Quality Inventory: 1996 Report to Congress, 36 percent of assessed river lengths and 39 percent of assessed lake areas were impaired for one or more of their designated uses (US EPA 1998). The most commonly reported causes of impairment in rivers and streams were siltation, nutrients, bacteria, oxygen-depleting substances, and pesticides; in lakes and reservoirs the causes also included metals and noxious aquatic plants. The most commonly reported sources of impairment were agriculture, nonpoint sources, municipal point sources, atmospheric deposition, hydrologic modification, habitat alteration and resource extraction. There were 2196 fish consumption advisories, which may include outright bans, in 47 States, the District of Columbia and American Samoa. Seventy-six percent of the advisories were due to mercury, with the rest due to PCBs, chlordane, dioxin, and DDT (US EPA 1998). States are not required to report fish kills for the National Inventory; however, available information for 1992 indicated 1620 incidents in 43 States, of which 930 were attributed to pollution, particularly oxygen-depleting substances, pesticides, manure, oil and gas, chlorine, and ammonia.

New approaches and tools, including appropriate technical guidance documents, are needed to facilitate ecosystem analyses of watersheds as required by the Clean Water Act. In particular, there is a pressing need for refinement and release of an ecological risk methodology that addresses the direct, indirect, and synergistic effects of nutrients, metals, toxic organic chemicals, and non-chemical stressors on aquatic ecosystems, including streams, rivers, lakes, and estuaries.

The ecosystem model AQUATOX is one of the few general ecological risk models that represents the combined environmental fate and effects of toxic chemicals. The model also represents conventional pollutants, such as nutrients and sediments, and considers several trophic levels, including attached and planktonic algae, submerged aquatic vegetation, several types of invertebrates, and several types of fish. It has been implemented for streams, small rivers, ponds, lakes, and reservoirs.

The AQUATOX model is described in these documents. **Volume 1: User's Manual** describes the usage of the model. Because the model is menu-driven and runs under Microsoft Windows on microcomputers, it is user-friendly and little guidance is required. **Volume 2: Technical Documentation** provides detailed documentation of the concepts and constructs of the model so that its suitability for given applications can be determined. **Volume 3: Validation Studies** presents three model validation studies performed for different environmental stressors and in different waterbody types.

1. INTRODUCTION

1.1 Overview

The AQUATOX model is a general ecological risk assessment model that represents the combined environmental fate and effects of conventional pollutants, such as nutrients and sediments, and toxic chemicals in aquatic ecosystems. It considers several trophic levels, including attached and planktonic algae and submerged aquatic vegetation, invertebrates, and forage, bottom-feeding, and game fish; it also represents associated organic toxicants (Figure 1). It can be implemented as a simple model (indeed, it has been used to simulate an abiotic flask) or as a truly complex food-web model. Often it is desirable to model a food web rather than a food chain, for example to examine the possibility of less tolerant organisms being replaced by more tolerant organisms as environmental perturbations occur. "Food web models provide a means for validation because they mechanistically describe the bioaccumulation process and ascribe causality to observed relationships between biota and sediment or water" (Connolly and Glaser 1998). The best way to accurately assess bioaccumulation is to use more complex models, but only if the data needs of the models can be met and there is sufficient time (Pelka 1998).

The model has been implemented for streams, small rivers, ponds, lakes, and reservoirs. The model is intended to be used to evaluate the likelihood of past, present, and future adverse effects from various stressors including potentially toxic organic chemicals, nutrients, organic wastes, sediments, and temperature. The stressors may be considered individually or together.

The fate portion of the model, which is applicable especially to organic toxicants, includes: partitioning among organisms, suspended and sedimented detritus, suspended and sedimented inorganic sediments, and water; volatilization; hydrolysis; photolysis; ionization; and microbial degradation. The effects portion of the model includes: chronic and acute toxicity to the various organisms modeled; and indirect effects such as release of grazing and predation pressure, increase in detritus and recycling of nutrients from killed organisms, dissolved oxygen sag due to increased decomposition, and loss of food base for animals.

AQUATOX represents the aquatic ecosystem by simulating the changing concentrations (in mg/L or g/m³) of organisms, nutrients, chemicals, and sediments in a unit volume of water (**Figure 1**). As such, it differs from population models, which represent the changes in numbers of individuals. As O'Neill et al. (1986) stated, ecosystem models and population models are complementary; one cannot take the place of the other. Population models excel at modeling individual species at risk and modeling fishing pressure and other age/size-specific aspects; but recycling of nutrients, the combined fate and effects of toxic chemicals, and other interdependencies in the aquatic ecosystem are important aspects that AQUATOX represents and that cannot be addressed by a population model.

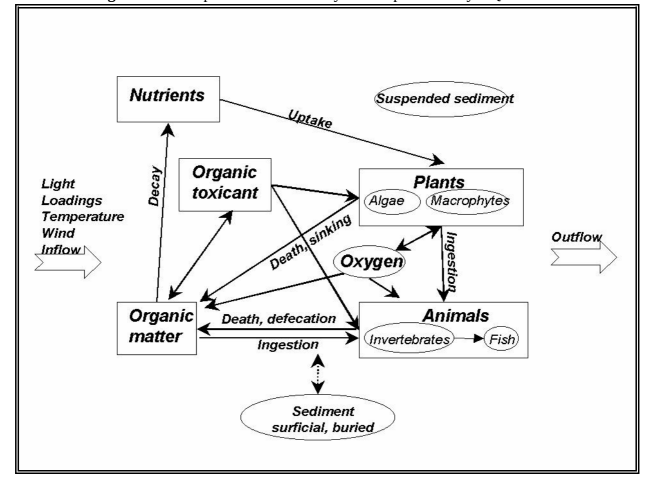


Figure 1. Conceptual Model of Ecosystem Represented by AQUATOX.

Any ecosystem model consists of multiple components requiring input data. These are the abiotic and biotic **state variables** or compartments being simulated (**Figure 2**). In AQUATOX the biotic state variables may represent trophic levels, guilds, and/or species. The model can represent a food web with both detrital- and algal-based trophic linkages. Closely related are **driving variables**, such as temperature, light, and nutrient loadings, which force the system to behave in certain ways. In AQUATOX state variables and driving variables are treated similarly in the code. This provides flexibility because external loadings of state variables, such as phytoplankton carried into a reach from upstream, may function as driving variables; and driving variables, such as pH and temperature, could be treated as dynamic state variables in a future implementation. Constant, dynamic, and multiplicative loadings can be specified for atmospheric, point- and nonpoint sources. Loadings of pollutants can be turned off at the click of a button to obtain a **control** simulation for comparison with the **perturbed** simulation.

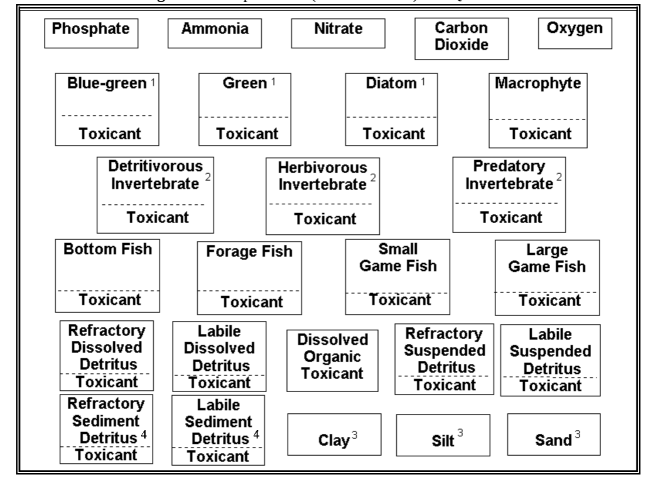


Figure 2. Compartments (State Variables) in AQUATOX

The model is written in object-oriented Pascal using the Delphi programming system for Windows. An object is a unit of computer code that can be duplicated; its characteristics and methods also can be inherited by higher-level objects. For example, the organism object, including variables such as the LC50 (lethal concentration of a toxicant) and process functions such as respiration, is inherited by the plant object; that is enhanced by plant-specific variables and functions and is duplicated for three kinds of algae; and the plant object is inherited and modified slightly for macrophytes. This modularity forms the basis for the remarkable flexibility of the model, including the ability to add and delete given state variables interactively.

AQUATOX utilizes **differential equations** to represent changing values of state variables, normally with a reporting time step of one day. These equations require starting values or **initial conditions** for the beginning of the simulation. If the first day of a simulation is changed, then the initial conditions may need to be changed. A simulation can begin with any date and may be for any length of time from a few days, corresponding to a microcosm experiment, to several years, corresponding to an extreme event followed by long-term recovery.

¹ Phytoplankton or Periphyton; ² Zooplankton or Zoobenthos;

³ Suspended, Sedimented, and Buried; ⁴ Surficial and buried

The **process equations** contain another class of input variables: the **parameters** or coefficients that allow the user to specify key process characteristics. For example, the maximum consumption rate is a critical parameter characterizing various consumers. AQUATOX is a mechanistic model with many parameters; however, default values are available so that the analyst only has to be concerned with those parameters necessary for a specific risk analysis, such as characterization of a new chemical. In the pages that follow, differential equations for the state variables will be followed by process equations and parameter definitions.

Finally, the system being modeled is characterized by **site constants**, such as mean and maximum depths. At present one can model small lakes, reservoirs, streams, small rivers, and ponds—and even enclosures and tanks. The generalized parameter screen is used for all these site types, although the hypolimnion entries obviously are not applicable to all. The temperature and light constants are used for simple forcing functions, blurring the distinctions between site constants and driving variables.

1.2 Background

AQUATOX is the latest in a long series of models, starting with the aquatic ecosystem model CLEAN (Park et al., 1974) and subsequently improved in consultation with numerous researchers at various European hydrobiological laboratories, resulting in the CLEANER series (Park et al., 1975, 1979, 1980; Park, 1978; Scavia and Park, 1976) and LAKETRACE (Collins and Park, 1989). The MACROPHYTE model, developed for the U.S. Army Corps of Engineers (Collins et al., 1985), provided additional capability for representing submersed aquatic vegetation. Another series started with the toxic fate model PEST, developed to complement CLEANER (Park et al., 1980, 1982), and continued with the TOXTRACE model (Park, 1984) and the spreadsheet equilibrium fugacity PART model. AQUATOX combined algorithms from these models with ecotoxicological constructs; and additional code was written as required for a truly integrative fate and effects model (Park, 1990, 1993). The model was then restructured and linked to Microsoft Windows interfaces to provide greater flexibility, capacity for additional compartments, and user friendliness (Park et al., 1995). The current version has been improved with the addition of constructs for chronic effects and uncertainty analysis, making it a powerful tool for probabilistic risk assessment (see Volume 3).

This technical documentation is intended to provide verification of individual constructs or mathematical and programming formulations used within AQUATOX. The scientific basis of the constructs reflects empirical and theoretical support; and precedence in the open literature and in widely used models is noted. Units are given to confirm the dimensional analysis. The mathematical formulations have been programmed and graphed in spreadsheets and the results have been evaluated in terms of behavior consistent with our understanding of ecosystem response; many of those graphs are given in the following documentation. The variable names in the documentation correspond to those used in the program so that the mathematical formulations and code can be compared, and the computer code has been checked for consistency with those formulations. Much of this has been done as part of the continuing process of internal review. This report is intended to expedite external review as well.

2. SIMULATION MODELING

2.1 Temporal and Spatial Resolution and Numerical Stability

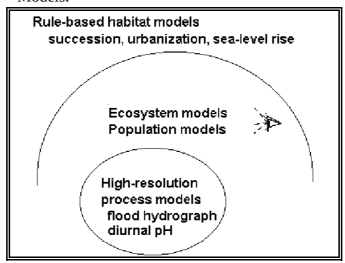
AQUATOX Release 1 is designed to be a general, realistic model of the fate and effects of pollutants in aquatic ecosystems. In order to be fast, easy to use, and verifiable, it has been designed with the simplest spatial and temporal resolutions consistent with this objective. It is designed to represent average daily conditions for a well-mixed aquatic system (in other words, a non-dimensional point model). It also can represent one-dimensional vertical epilimnetic and hypolimnetic conditions for those systems that exhibit stratification on a seasonal basis.

According to Ford and Thornton (1979), a one-dimensional model is appropriate for reservoirs that are between 0.5 and 10 km in length; if larger, then a two-dimensional model disaggregated along the long axis is indicated. The one-dimensional assumption is also appropriate for many lakes (Stefan and Fang, 1994). Similarly, one can consider a single reach or stretch of river at a time. A distributed version of the model (Version 2.00) is being developed; it will be able to simulate several linked stream reaches.

Usually the reporting time step is one day, but numerical instability is avoided by allowing the step size of the integration to vary to achieve a predetermined accuracy in the solution. This is a numerical approach, and the step size is not directly related to the temporal scale of the ecosystem simulation. AQUATOX uses a very efficient fourth- and fifth-order Runge-Kutta integration routine with adaptive step size to solve the differential equations (Press et al., 1986, 1992). The routine uses the fifth-order solution to determine the error associated with the fourth-order solution; it decreases the step size (often to 15 minutes or less) when rapid changes occur and increases the step size when there are slow changes, such as in winter. However, the step size is constrained to a maximum of one day so that short-term pollutant loadings are always detected.

The temporal and spatial resolution is in keeping with the generality and realism of the model (see Park and Collins, 1982). Careful consideration has been given to the hierarchical nature of the system. Hierarchy theory tells us that models should have resolutions appropriate to the objectives; phenomena with temporal and spatial scales that are significantly longer than those of interest should be treated as constants, and phenomena with much smaller temporal and spatial scales should be treated as steady-state properties or parameters (**Figure 3**, O'Neill et al., 1986). The model uses a longer time step than dynamic hydrologic models that are concerned with representing short-term phenomena such as storm hydrographs, and it uses a shorter time step than fate models that may be concerned only with long-term patterns such as bioaccumulation in large fish.

Figure 3. Position of Ecosystem Models such as AQUATOX in the Spatial-temporal Hierarchy of Models.



Changing the permissible relative error (the difference between the fourth- and fifth-order solutions) of the simulation can affect the results. The model allows the user to set the relative error, usually between 0.005 and 0.01. Comparison of output shows that up to a point a smaller error can yield a marked improvement in the simulation—although execution time is slightly longer. For example, simulations of two pulsed doses of chlorpyrifos in a pond exhibit a spread in the first pulse of about $0.6~\mu g/L$ dissolved toxicant between the simulation with 0.001 relative error and the simulation with 0.05 relative error (**Figure 4**); this is probably due in part to differences in the timing of the reporting step. However, if we examine the dissolved oxygen levels, which combine the effects of photosynthesis, decomposition, and reaeration, we find that there are pronounced differences over the entire simulation period. The simulations with 0.001 and 0.01 relative error give almost exactly the same results, suggesting that the more efficient 0.001 relative error should be used; the simulation with 0.05 relative error exhibits instability in the oxygen simulation; and the simulation with 0.1 error gives quite different values for dissolved oxygen (**Figure 5**). The observed mean daily maximum dissolved oxygen for that period was 9.2 mg/L (US EPA 1988), which corresponds most closely with the results of simulation with 0.001 and 0.01 relative error.

Figure 4. Pond with Chlorpyrifos in Dissolved Phase.

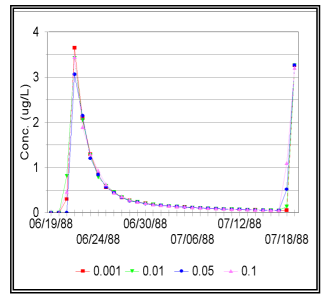
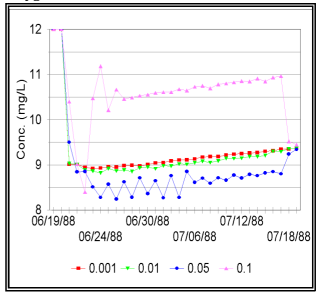


Figure 5. Same as **Figure 4** with Dissolved Oxygen.



2.2 Uncertainty Analysis

There are numerous sources of uncertainty and variation in natural systems. These include: site characteristics such as water depth, which may vary seasonally and from site to site; environmental loadings such as water flow, temperature, and light, which may have a stochastic component; and critical biotic parameters such as maximum photosynthetic and consumption rates, which vary among experiments and representative organisms.

In addition, there are sources of uncertainty and variation with regard to pollutants, including: pollutant loadings from runoff, point sources, and atmospheric deposition, which may vary stochastically from day to day and year to year; physico-chemical characteristics such as octanol-water partition coefficients and Henry Law constants that cannot be measured easily; chemodynamic parameters such as microbial degradation, photolysis, and hydrolysis rates, which may be subject to both measurement errors and indeterminate environmental controls.

Increasingly, environmental analysts and decision makers are requiring probabilistic modeling approaches so that they can consider the implications of uncertainty in the analyses. AQUATOX provides this capability by allowing the user to specify the types of distribution and key statistics for a wide selection of input variables. Depending on the specific variable and the amount of available information, any one of several distributions may be most appropriate. A lognormal distribution is the default for environmental and pollutant loadings. In the uncertainty analysis, the distributions for constant loadings are sampled daily, providing day-to-day variation within the limits of the distribution, reflecting the stochastic nature of such loadings. Distributions for dynamic loadings may employ multiplicative factors that are sampled once each simulation (**Figure 6**). Normally the multiplicative factor for a loading is set to 1, but, as seen in the example, under

extreme conditions the loading may be ten times as great. In this way the user could represent unexpected conditions such as pesticides being applied inadvertently just before each large storm of the season. Loadings usually exhibit a lognormal distribution, and that is suggested in these applications, unless there is information to the contrary.

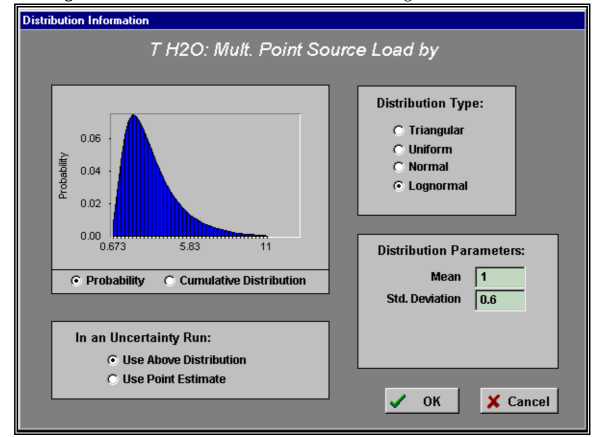


Figure 6. Distribution Screen for Point-Source Loading of Toxicant in Water.

A sequence of increasingly informative distributions should be considered for most parameters (see **Volume 1: User's Manual**.) If only two values are known and nothing more can be assumed, the two values may be used as minimum and maximum values for a uniform distribution (**Figure 7**); this is often used for parameters where only two values are known. If minimal information is available but there is reason to accept a particular value as most likely, perhaps based on calibration, then a triangular distribution may be most suitable (**Figure 8**). Note that the minimum and maximum values for the distribution are constraints that have zero probability of occurrence. If additional data are available indicating both a central tendency and spread of response, such as parameters for well-studied processes, then a normal distribution may be most appropriate (**Figure 9**). The result of applying such a distribution in a simulation of Onondaga Lake, New York is shown in **Figure 10**, where simulated benthic feeding is seen to affect the sediment-water interaction and subsequently the predicted hypolimnetic anoxia. All distributions are truncated at zero because negative values would have no meaning. A non-random seed can be used for the

random number generator, causing the same sequence of numbers to be picked in successive applications; this is useful if you want to be able to duplicate the results exactly.

Figure 7. Uniform Distribution for Henry's Law Constant for Esfenvalerate.

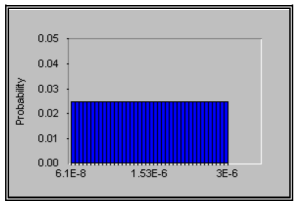


Figure 8. Triangular Distribution for Maximum Consumption Rate for Bass.

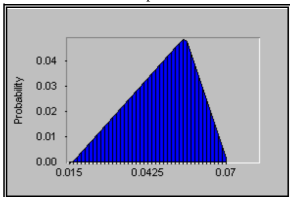
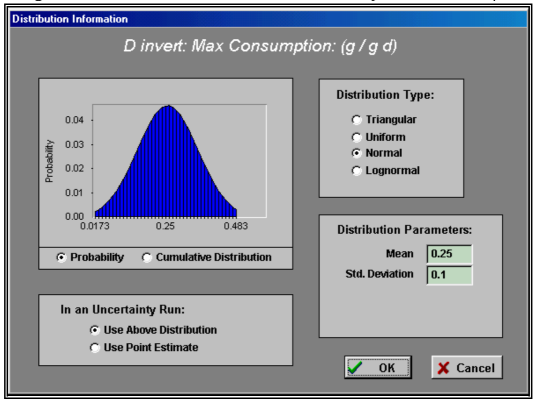


Figure 9. Normal Distribution for Maximum Consumption Rate for *Tubifex*.



Efficient sampling from the distributions is obtained with the Latin hypercube method (McKay et al., 1979; Palisade Corporation, 1991), using algorithms originally written in FORTRAN (Anonymous, 1988). Depending on how many iterations are chosen for the analysis, each cumulative distribution is subdivided into that many equal segments. Then a uniform random value is chosen *within* each segment and used in one of the subsequent simulation runs. For example, the

distribution shown in **Figure 9** can be sampled as shown in **Figure 11**. This method is particularly advantageous because all regions of the distribution, including the tails, are sampled. The default is twenty iterations, meaning that twenty simulations will be performed with sampled input values; this should be considered the minimum number to provide any reliability. The optimal number can be determined experimentally by noting the number required to obtain convergence of mean response values for key state variables; in other words, at what point do additional iterations not result in significant changes in the results? As many variables may be represented by distributions as desired, but the method assumes that they are independently distributed. By varying one parameter at a time the sensitivity of the model to individual parameters can be determined. This is done for key parameters in the following documentation.

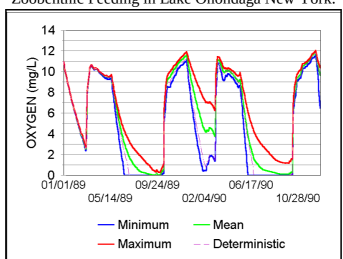
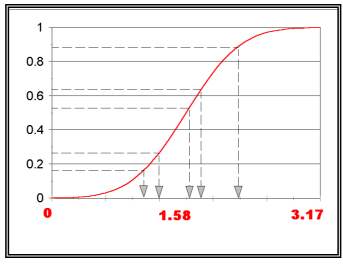


Figure 10. Sensitivity of Hypolimnetic Oxygen to Zoobenthic Feeding in Lake Onondaga New York.

Figure 11. Latin Hypercube Sampling of a Cumulative Distribution with a Mean of 25 and Standard Deviation of 8 Divided into 5 Intervals.



3. PHYSICAL CHARACTERISTICS

3.1 Morphometry

Volume

Volume is a state variable and can be computed in several ways depending on availability of data and the site dynamics. It is important for computing the dilution or concentration of pollutants, nutrients, and organisms; it may be constant, but usually it is time varying. In the model, ponds, lakes, and reservoirs are treated differently than streams, especially with respect to computing volumes. The change in volume of ponds, lakes, and reservoirs is computed as:

$$\frac{dVolume}{dt} = Inflow - Discharge - Evap$$
 (1)

where:

dVolume/dt = derivative for volume of water (m³/d), Inflow = inflow of water into waterbody (m³/d),

Discharge = discharge of water from waterbody (m^3/d), and

Evap = evaporation (m^3/d) , see (2).

Evaporation is converted from an annual value for the site to a daily value using the simple relationship:

$$Evap = \frac{MeanEvap}{365} \cdot 0.0254 \cdot Area \tag{2}$$

where:

MeanEvap = mean annual evaporation (in/yr),

= days per year (yr),

0.0254 = conversion from inches to meters (m/in), and

Area = area of the waterbody (m²).

The user is given several options for computing volume including keeping the volume constant; making the volume a dynamic function of inflow, discharge, and evaporation; using a time series of known values; and computing volume as a function of the Manning's equation. Depending on the method, inflow and discharge are varied, as indicated in **Table 1**.

Table 1. Computation of Volume, Inflow, and Discharge

Method	Inflow	Discharge	
Constant	InflowLoad	InflowLoad - Evap	
Dynamic	InflowLoad	DischargeLoad	
Known values	InflowLoad	InflowLoad - Evap + (State - KnownVals)/dt	
Manning	ManningVol - (State + Discharge)/dt + Evap	DischargeLoad	

The variables are defined as:

InflowLoad = user-supplied inflow loading (m³/d); DischargeLoad = user-supplied discharge loading (m³/d);

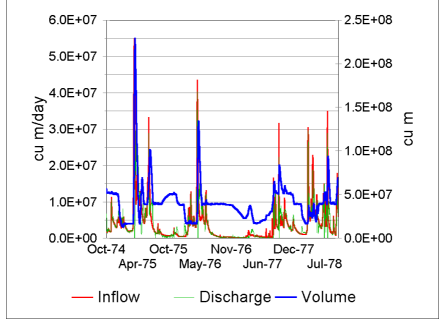
State = computed state variable value for volume (m³); KnownVals = time series of known values of volume (m³); dt = incremental time in simulation (d); and ManningVol = volume of stream reach (m³), see (3).

Figure 12 illustrates time-varying volumes and inflow loadings specified by the user and discharge computed by the model for a run-of-the-river reservoir. Note that significant drops in volume occur with operational releases, usually in the spring, for flood control purposes.

Figure 12. Volume, Inflow, and Discharge for a 4-year Period in Coralville Reservoir, Iowa.

6.0E+07

2.5E+08



The time-varying volume of water in a stream channel is computed as:

$$ManningVol = Y \cdot CLength \cdot Width$$
 (3)

where:

Y = dynamic mean depth (m), see (4);

CLength = length of reach (m); and Width = width of channel (m).

In streams the depth of water and flow rate are key variables in computing the transport, scour, and deposition of sediments. Time-varying water depth is a function of the flow rate, channel roughness, slope, and channel width using Manning's equation:

$$Y = \left(\frac{Q \cdot Manning}{\sqrt{Slope} \cdot Width}\right)^{3/5} \tag{4}$$

where:

Q = flow rate (m³/s);

Manning = Manning's roughness coefficient $(s/m^{1/3})$;

Slope = slope of channel (m/m); and

Width = channel width (m).

The Manning's roughness coefficient is an important parameter representing frictional loss, but it is not subject to direct measurement. The user can choose among the following stream types:

- concrete channel (with a default Manning's coefficient of 0.020);
- dredged channel, such as ditches and channelized streams (default coefficient of 0.030); and
- natural channel (default coefficient of 0.040).

These generalities are based on Chow's (1959) tabulated values as given by Hoggan (1989).

In the absence of inflow data, the flow rate is computed from the initial mean water depth, assuming a rectangular channel and using a rearrangement of Manning's equation:

$$QBase = \frac{IDepth^{5/3} \cdot \sqrt{Slope} \cdot Width}{Manning}$$
 (5)

where:

QBase = base flow (m^3/s) ; and

Idepth = mean depth as given in site record (m).

The dynamic flow rate is calculated from the inflow loading by converting from m³/d to m³/s:

$$Q = \frac{Inflow}{86400} \tag{6}$$

where:

Q = flow rate (m³/s); and

Inflow = water discharged into channel from upstream (m^3/d) .

Bathymetric Approximations

The depth distribution of a water body is important because it determines the areas and volumes subject to mixing and light penetration. The shapes of ponds, lakes, reservoirs, and streams are represented in the model by idealized geometrical approximations, following the topological

treatment of Junge (1966; see also Straškraba and Gnauck, 1985). The shape parameter *P* (Junge, 1966) characterizes the site, with a shape that is indicated by the ratio of mean to maximum depth.:

$$P = 6.0 \cdot \frac{ZMean}{ZMax} - 3.0 \tag{7}$$

Where:

ZMean = mean depth (m);

ZMax = maximum depth (m); and

P = characterizing parameter for shape (unitless).; P is constrained

between -1.0 and 1.0

Shallow constructed ponds and ditches may be approximated by an ellipsoid where Z/ZMax = 0.6 and P = 0.6. Reservoirs generally are extreme elliptic sinusoids with values of P constrained to -1.0. Lakes may be either elliptic sinusoids, with P between 0.0 and -1.0, or elliptic hyperboloids with P between 0.0 and 1.0 (**Table 2**). The model requires mean and maximum depth, but if only the maximum depth is known, then the mean depth can be estimated by multiplying ZMax by the representative ratio. Not all water bodies fit the elliptic shapes, but the model generally is not sensitive to the deviations.

Based on these relationships, fractions of volumes and areas can be determined for any given depth (Junge, 1966) (**Figure 13-Figure 14**):

$$AreaFrac = (1.0 + P) \cdot \frac{Z}{ZMax} - P \cdot (\frac{Z}{ZMax})^{2}$$
 (8)

$$VolFrac = \frac{6.0 \cdot \frac{Z}{ZMax} - 3.0 \cdot (1.0 - P) \cdot (\frac{Z}{ZMax})^{2} - 2.0 \cdot P \cdot (\frac{Z}{ZMax})^{3}}{3.0 + P}$$
(9)

where:

AreaFrac = fraction of area of site above given depth (unitless);

VolFrac = fraction of volume of site above given depth (unitless); and

Z = depth of interest (m).

Table 2. Examples of Morphometry of Waterbodies

Table 2. Examples	of Morphome	try of wa	terboules
Site	ZMean/ZMax	Р	Constrained P
Lakes			
Chad, Chad	0.13	-2.22	-1.00
Managua, Nicaragua	0.26	-1.42	-1.00
Michigan, U.SCanada	0.27	-1.38	-1.00
Erie, U.SCanada	0.33	-1.02	-1.00
Windermere, England	0.36	-0.85	-0.85
Baikal, Russia	0.43	-0.42	-0.42
Como, Italy	0.45	-0.30	-0.30
Superior, U.SCanada	0.47	-0.18	-0.18
Tahoe, CA-NV	0.50	0.00	0.00
Esrom, Denmark	0.56	0.35	0.35
Clear, CA	0.57	0.43	0.43
Crater, OR	0.60	0.60	0.60
Kinneret, Israel	0.60	0.63	0.63
Okeechobee, FL	0.67	1.00	1.00
Ontario, U.SCanada	0.69	1.14	1.00
Balaton, Hungary	0.75	1.50	1.00
George, Uganda	0.80	1.80	1.00
Reservoirs			
DeGray, AR	0.25	-1.49	-1.00
Grenada, MS	0.21	-1.74	-1.00
Lewis and Clark, SD	0.31	-1.13	-1.00
Texoma, TX	0.27	-1.38	-1.00
Delaware, OH	0.22	-1.68	-1.00
Sidney Lanier, GA	0.33	-1.01	-1.00
Monroe, IN	0.30	-1.18	-1.00
Tenkiller Ferry, OK	0.36	-0.86	-0.86
Mendocino, CA	0.36	-0.84	-0.84
Coralville, IA	0.37	-0.80	-0.80
Waterbury, VT	0.43	-0.42	-0.42
Pend Oreille, ID	0.50	-0.03	-0.03
Ponds			
Czech Rep., fish (very old)	0.43	-0.42	-0.42
Czech Rep., Elbe R. backwaters	0.50	-0.03	-0.03
Dor. Israel, fish, recent	0.67	1.00	1.00

data from Hutchinson, 1957; HrbáČek, 1966; Leidy and Jenkins, 1977; and Horne and Goldman, 1994

For example, the fraction of the volume that is epilimnion can be computed by setting depth Z to the mixing depth. Furthermore, by setting Z to the depth of the euphotic zone, the fraction of the fraction of the area available for colonization by macrophytes and periphyton can be computed:

FracLit =
$$(1 + P) \cdot \frac{ZEuphotic}{ZMax} - P \cdot \left(\frac{ZEuphotic}{ZMax}\right)^2$$
 (10)

If the site is a limnocorral (an artificial enclosure) then the available area is increased accordingly:

$$FracLittoral = FracLit \cdot \frac{Area + LimnoWallArea}{Area}$$
 otherwise (11)

FracLittoral = FracLit

where:

FracLittoral fraction of site area that is within the euphotic zone (unitless); **Z**Euphotic depth of the euphotic zone, where primary production exceeds

respiration, usually calculated as a function of extinction (m);

site area (m²); and Area =

area of limnocorral walls (m²). LimnoWallArea

The depth of the euphotic zone, where radiation is 1% of surface radiation, is computed as (Thomann and Mueller, 1987):

$$ZEuphotic = 4.605/Extinct$$
 (12)

where:

Extinct the overall extinction coefficient (1/m), see (30).

Figure 13 Volume as a Function of Depth in Ponds

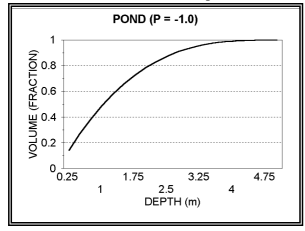
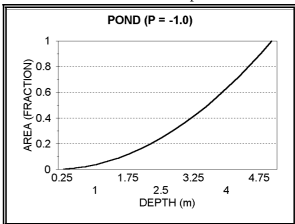


Figure 14 Area as a Function of Depth in Ponds



3.2 Washout

Transport out of the system, or washout, is an important loss term for nutrients, floating organisms, and dissolved toxicants in reservoirs and streams. Although it is considered separately for several state variables, the process is a general function of discharge:

$$Washout = \frac{Discharge}{Volume} \cdot State$$
 (13)

where:

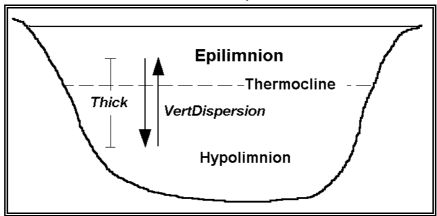
Washout = loss due to being carried downstream ($g/m^3 \cdot d$), and

State = concentration of dissolved or floating state variable (g/m^3) .

3.3 Stratification and Mixing

Thermal stratification is handled in the simplest form consistent with the goals of forecasting the effects of nutrients and toxicants. Lakes and reservoirs are considered in the model to have two vertical zones: epilimnion and hypolimnion (**Figure 15**); the metalimnion zone that separates these is ignored. Instead, the thermocline, or plane of maximum temperature change, is taken as the separator; this is also known as the mixing depth (Hanna, 1990). Dividing the lake into two vertical zones follows the treatment of Imboden (1973), Park et al. (1974), and Straškraba and Gnauck (1983). The onset of stratification is considered to occur when the mean water temperature exceeds 4° and the difference in temperature between the epilimnion and hypolimnion exceeds 3°. Overturn occurs when the temperature of the epilimnion is less than 3°, usually in the fall. Winter stratification is not modeled. For simplicity, the thermocline is assumed to occur at a constant depth.

Figure 15
Thermal Stratification in a Lake; Terms Defined in Text



There are numerous empirical models relating thermocline depth to lake characteristics. AQUATOX uses an equation by Hanna (1990), based on the maximum effective length (or fetch). The dataset includes 167 mostly temperate lakes with maximum effective lengths of 172 to 108,000 m and ranging in altitude from 10 to 1897 m. The equation has a coefficient of determination $r^2 =$

0.850, meaning that 85 percent of the sum of squares is explained by the regression. Its curvilinear nature is shown in **Figure 16**, and it is computed as (Hanna, 1990):

$$\log(MaxZMix) = 0.336 \cdot \log(Length) - 0.245 \tag{14}$$

where:

MaxZMix = maximum mixing depth for lake (m); and Length = maximum effective length for wave setup (m).

Wind action is implicit in this formulation. Wind has been modeled explicitly by Baca and Arnett (1976, quoted by Bowie et al., 1985), but their approach requires calibration to individual sites, and it is not used here.

Vertical dispersion for bulk mixing is modeled as a function of the time-varying hypolimnetic and epilimnetic temperatures, following the treatment of Thomann and Mueller (1987, p. 203; see also Chapra and Reckhow, 1983, p. 152; **Figure 17**):

$$VertDispersion = Thick \cdot \left(\frac{HypVolume}{ThermoclArea \cdot Deltat} \cdot \frac{|T_{hypo}^{t-1} - T_{hypo}^{t+1}|}{T_{epi}^{t} - T_{hypo}^{t}} \right)$$
(15)

where:

VertDispersion = vertical dispersion coefficient (m²/d);

Thick = distance between the centroid of the epilimnion and the centroid of the

hypolimnion, effectively the mean depth (m);

HypVolume = volume of the hypolimnion (m³); ThermoclArea = area of the thermocline (m²);

Deltat = time step (d);

 T_{hypo}^{t-1} , T_{hypo}^{t+1} = temperature of hypolimnion one time step before and one time step

after present time (°C); and

 T_{epi}^{t} , T_{hypo}^{t} = temperature of epilimnion and hypolimnion at present time (°C).

MAXIMUM MIXING DEPTH

25
20
100
11500
22900
34300
5800
17200
28600
40000

Figure 16Mixing Depth as a Function of Fetch

Stratification can break down temporarily as a result of high throughflow. This is represented in the model by making the vertical dispersion coefficient between the layers a function of discharge for sites with retention times of less than or equal to 180 days (**Figure 18**), rather than temperature differences as in equation **15**, based on observations by Straškraba (1973) for a Czech reservoir:

LENGTH (m)

$$VertDispersion = 1.37 \cdot 10^4 \cdot Retention^{-2.269}$$
 (16)

and:

$$Retention = \frac{Volume}{TotDischarge}$$
 (17)

where:

Retention = retention time (d); $Volume = volume of site (m^3); and$ $TotDischarge = total discharge (m^3/d).$

Figure 17
Vertical Dispersion as a Function of Temperature Differences

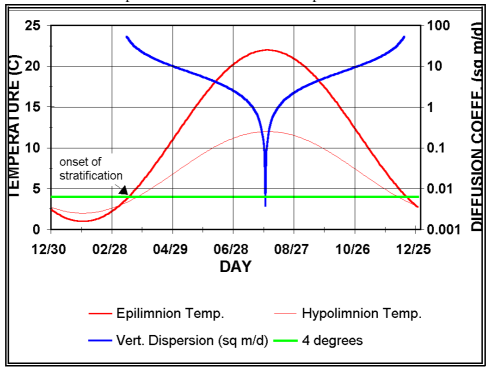
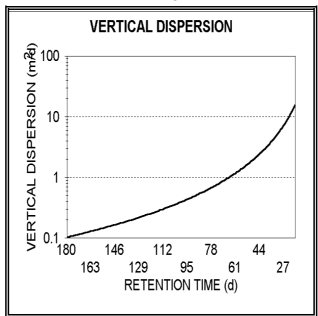


Figure 18
Vertical Dispersion as a Function of Retention
Time



The bulk vertical mixing coefficient is computed using site characteristics and the time-varying vertical dispersion (Thomann and Mueller, 1987):

$$BulkMixCoeff = \frac{VertDispersion \cdot ThermoclArea}{Thick}$$
 (18)

where:

BulkMixCoeff = bulk vertical mixing coefficient (m 3 /d),

ThermoclArea = area of thermocline (m^2) .

Turbulent diffusion between epilimnion and hypolimnion is computed separately for each segment for each time step while there is stratification:

$$TurbDiff_{epi} = \frac{BulkMixCoeff}{Volume_{epi}} \cdot (Conc_{compartment, hypo} - Conc_{compartment, epi})$$
 (19)

$$TurbDiff_{hypo} = \frac{BulkMixCoeff}{Volume_{hypo}} \cdot (Conc_{compartment, epi} - Conc_{compartment, hypo})$$
(20)

where:

TurbDiff = turbulent diffusion for a given zone (g/m³·d);

Volume = volume of given segment (m^3) ; and

Conc = concentration of given compartment in given zone (g/m^3) .

The effects of stratification, mixing due to high throughflow, and overturn are well illustrated by the pattern of dissolved oxygen levels in the hypolimnion of Lake Nockamixon, a eutrophic reservoir in Pennsylvania (**Figure 19**).

Pennsylvania as Shown by Hypolimnetic Dissolved Oxygen

14

(1)
12
onset of stratification
whigh throughflow
overturn
stratification
high throughflow

Figure 19
Stratification and Mixing in Lake Nockamixon,
Pennsylvania as Shown by Hypolimnetic Dissolved Oxygen

3.4 Temperature

Default water temperature loadings for the epilimnion and hypolimnion are represented through a simple sine approximation for seasonal variations (Ward, 1963) based on user-supplied observed means and ranges (**Figure 20**):

01/01/82 03/07/82 05/11/82 07/15/82 09/18/82 11/22/82

Temperature = TempMean +
$$(-1.0 \cdot \frac{TempRange}{2} \cdot (\sin(0.0174533 \cdot (0.987 \cdot (Day + PhaseShift) - 30))))]$$
 (21)

where:

Temperature = average daily water temperature (°C);

TempMean = mean annual temperature (°C);TempRange = annual temperature range (°C),

Day = Julian date (d); and

PhaseShift = time lag in heating (= 90 d).

Observed temperature loadings should be entered if responses to short-term variations are of interest. This is especially important if the timing of the onset of stratification is critical, because stratification is a function of the difference in hypolimnetic and epilimnetic temperatures (see **Figure 18**).

3.5 Light

The default incident light function is a variation on the temperature equation, but without the lag term:

$$Solar = LightMean + \frac{LightRange}{2} \cdot \sin(0.0174533 \cdot Day - 1.76)$$
 (22)

where:

Solar = average daily incident light intensity (ly/d);

LightMean = mean annual light intensity (ly/d);

LightRange = annual range in light intensity (ly/d); and

Day = Julian date (d).

The derived values are given as average light intensity in Langleys per day (Ly/d = 10 kcal/m²·d). An observed time-series of light also can be supplied by the user; this is especially important if the effects of daily climatic conditions are of interest. If the average water temperature drops below 3°C, the model assumes the presence of ice cover and decreases light to 33% of incident radiation. This reduction, due to the reflectivity and transmissivity of ice and snow, is an average of widely varying values summarized by Wetzel (1975; also see LeCren and Lowe-McConnell, 1980). The model does not automatically adjust for shading by riparian vegetation, so a times-series should probably be supplied if modeling a narrow stream.

Photoperiod is approximated using the Julian date (**Figure 21**):

$$Photoperiod = \frac{12 + A \cdot \cos(380 \cdot \frac{Day}{365} + 248)}{24}$$
 (23)

where:

Photoperiod = fraction of the day with daylight (unitless);

A = hours of daylight minus 12 (hr); and

Day = Julian date (d).

A is the difference between the number of hours of daylight at the summer solstice at a given latitude and the vernal equinox, and is given by a linear regression developed by Groden (1977):

$$A = 0.1414 \cdot Latitude - Sign \cdot 2.413 \tag{24}$$

where:

Latitude = latitude ($^{\circ}$, decimal), negative in southern hemisphere; and Sign = 1.0 in northern hemisphere, -1.0 in southern hemisphere.

Figure 20Annual Temperature

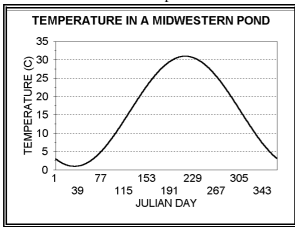
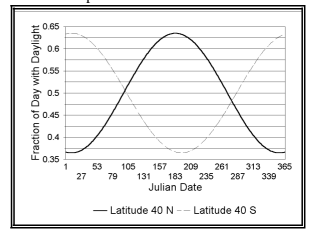


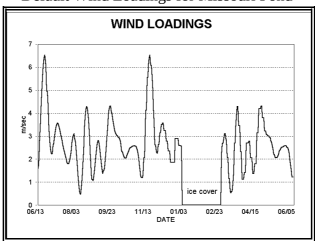
Figure 21
Photoperiod as a Function of Date



3.6 Wind

Wind is an important driving variable because it determines the stability of blue-green algal blooms, and reaeration or oxygen exchange, and it controls volatilization of some organic chemicals. If site data are not available, default variable wind speeds are represented through a Fourier series of sine and cosine terms; the mean and first ten harmonics seem to capture the variation adequately (**Figure 22**). This default loading is based on an unpublished 140-day record (May 20 to October 12) from Columbia, Missouri; therefore, it has a 140-day repeat, representative of the Midwest during the growing season. This approach is quite useful because the mean can be specified by the user and the variability will be imposed by the function. If ice cover is predicted, wind is set to 0.

Figure 22
Default Wind Loadings for Missouri Pond



4. BIOTA

The biota consists of two main groups, plants and animals; each is represented by a set of process-level equations. In turn, plants are differentiated into algae and macrophytes, represented by slight variations in the differential equations. Algae may be either phytoplankton or periphyton. Phytoplankton are subject to sinking and washout, while periphyton are subject to substrate limitation and scour by currents. These are treated as process-level differences in the equations.

Animals are subdivided into invertebrates and fish; the invertebrates may be pelagic invertebrates, benthic insects or other benthic invertebrates. These groups are represented by different parameter values and by variations in the equations. Insects are subject to emergence, but benthic invertebrates are not. Gamefish may be represented by both young of the year and adults, which are connected by promotion.

4.1 Algae

The change in algal biomass—expressed as g/m³ for phytoplankton, but as g/m² for periphyton—is a function of the loading (especially phytoplankton from upstream), photosynthesis, respiration, excretion or photorespiration, nonpredatory mortality, grazing or predatory mortality, and washout; as noted above, phytoplankton also are subject to sinking. If the system is stratified, turbulent diffusion also affects the biomass of phytoplankton:

$$\frac{dBiomass}{dt} = Loading + Photosynthesis - Respiration - Excretion - Mortality - Predation \pm Sinking - Washout \pm TurbDiff$$
 (25)

where:

dBiomass/dt = change in biomass of algae with respect to time (g/m³·d);

Loading = loading of algal group $(g/m^3 \cdot d)$;

Photosynthesis = rate of photosynthesis (g/m^3 ·d), see (26);

Respiration = respiratory loss ($g/m^3 \cdot d$), see (51);

Excretion = excretion or photorespiration ($g/m^3 \cdot d$), see (52);

Mortality = nonpredatory mortality ($g/m^3 \cdot d$), see (54);

Predation = herbivory $(g/m^3 \cdot d)$, see (74);

Washout = loss due to being carried downstream ($g/m^3 \cdot d$), see (60);

Sinking = loss or gain due to sinking between layers and sedimentation to bottom

 $(g/m^3 \cdot d)$, see (57); and

TurbDiff = turbulent diffusion (g/m 3 ·d), see (18).

Figure 23 and Figure 24 are examples of the predicted changes in biomass and the processes that contribute to these changes in a eutrophic lake.

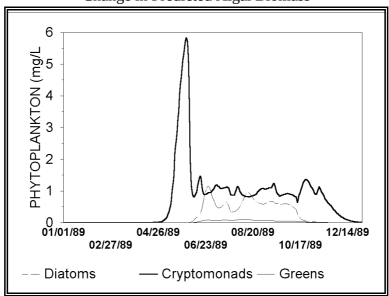
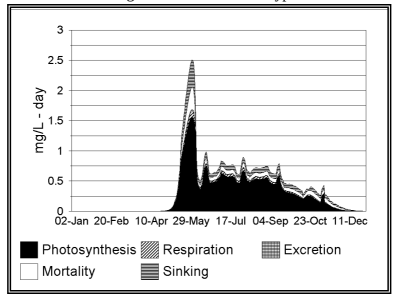


Figure 23
Change in Predicted Algal Biomass

Figure 24
Predicted Algal Process Rates in Cryptomonads



Photosynthesis is modeled as a maximum observed rate multiplied by reduction factors for the effects of toxicants and suboptimal light, temperature, current, and nutrients:

$$Photosynthesis = PMax \cdot PProdLimit \cdot Biomass$$
 (26)

The limitation of primary production in phytoplankton is:

$$PProdLimit = LtLimit \cdot NutrLimit \cdot VLimit \cdot TCorr \cdot FracPhoto$$
 (27)

Periphyton have an additional limitation based on available substrate:

$$PProdLimit = LtLimit \cdot NutrLimit \cdot VLimit \cdot TCorr \cdot FracPhoto \cdot FracLittoral$$
 (28)

where:

Pmax = maximum photosynthetic rate (1/d); LtLimit = light limitation (unitless), see (29); NutrLimit = nutrient limitation (unitless), see (43);

Vlimit = current limitation for periphyton (unitless), see (44);

TCorr = limitation due to suboptimal temperature (unitless), see (47);

FracPhoto = reduction factor for effect of toxicant on photosynthesis (unitless), see

(271);

FracLittoral = fraction of area that is within euphotic zone (unitless) see (11); and

Biomass = biomass of algae (g/m^3) .

Under optimal conditions, a reduction factor has a value of 1; otherwise, it has a fractional value. Use of a multiplicative construct implies that the factors are independent. Several authors (for example, Collins, 1980; Straškraba and Gnauck, 1983) have shown that there are interactions among the factors. However, we feel the data are insufficient to generalize to all algae; therefore, the simpler multiplicative construct is used, as in many other models (Chen and Orlob, 1975; Lehman et al., 1975; Jørgensen, 1976; DiToro et al., 1977; Kremer and Nixon, 1978; Park et al., 1985; Ambrose et al., 1991). Default parameter values for the various processes are taken primarily from compilations (for example, Jørgensen, 1979; Collins and Wlosinski, 1983; Bowie et al., 1985); they may be modified as needed.

Light Limitation

Because it is required for photosynthesis, light is a very important limiting variable. It is especially important in controlling competition among plants with differing light requirements. Similar to many other models (for example, Di Toro et al., 1971; Park et al., 1974, 1975, 1979, 1980; Lehman et al., 1975; Canale et al., 1975, 1976; Thomann et al., 1975, 1979; Scavia et al., 1976; Bierman et al., 1980; O'Connor et al., 1981), AQUATOX uses the Steele (1962) formulation for light limitation. Light is specified as average daily radiation. The average radiation is multiplied by the photoperiod, or the fraction of the day with sunlight, based on a simplification of Steele's (1962) equation proposed by Di Toro et al. (1971):

$$LtLimit = 0.85 \cdot \frac{e}{Extinct \cdot (DepthBottom - DepthTop)}$$

$$\cdot Photoperiod \cdot (LtAtDepth - LtAtTop)$$
(29)

where:

LtLimit = light limitation (unitless);

e = the base of natural logarithms (2.71828, unitless);

Photoperiod = fraction of day with daylight, see (23); Extinct = total light extinction (1/m), see (30); DepthBottom = maximum depth or depth of bottom of layer if stratified (m); if

periphyton or macrophyte then limited to euphotic depth;

DepthTop = depth of top of layer (m);

LtAtDepth = see (32); and LtAtTop = see (33), (34).

Because the equation overestimates by 15 percent the cumulative effect of light limitation over a 24-hour day, a correction factor of 0.85 is applied (Kremer and Nixon, 1978).

Extinction of light is based on several additive terms: the baseline extinction coefficient for pure water, the so-called "self-shading" of plants, attenuation due to suspended particulate organic matter (POM) and inorganic sediment, and attenuation due to dissolved organic matter (DOM):

$$Extinct = WaterExtinction + PhytoExtinction + ECoeffDOM \cdot DOM + ECoeffPOM \cdot \Sigma PartDetr + ECoeffSed \cdot InorgSed$$
(30)

where:

WaterExtinction = extinction due to pure water (1/m);

PhytoExtinction = extinction due to phytoplankton and periphyton (1/m), see (31);

EcoeffDOM = attenuation coefficient for dissolved detritus (1/m-g/m³);

DOM = concentration of dissolved organic matter (g/m³), see (96) and (97);

EcoeffPOM = attenuation coefficient for particulate detritus (1/m-g/m³); PartDetr = concentration of particulate detritus (g/m³), see (94) and (95); EcoeffSed = attenuation coefficient for suspended sediment (1/m-g/m³); and

InorgSed = total suspended inorganic sediment (g/m³), see (177).

For computational reasons, the value of *Extinct* is constrained between 5⁻¹⁹ and 25. Self-shading by phytoplankton, periphyton, and macrophytes is a function of the biomass and attenuation coefficient for each group:

$$PhytoExtinction = \sum_{alga} (ECoeffPhyto_{alga} \cdot Biomass_{alga})$$
(31)

where:

EcoeffPhyto = attenuation coefficient for given alga (1/m-g/m³); and

Biomass = concentration of given alga (g/m^3) .

The light at depth is computed by:

$$LtAtDepth = e^{-\frac{Light}{LightSat}} \cdot e^{-Extinction \cdot DepthBottom}$$
(32)

Light at the surface of the waterbody is computed by:

$$LtAtTop = e^{-\frac{Light}{LightSat}}$$
 (33)

and light at the top of the hypolimnion is computed by:

$$LtAtTop = e^{-\frac{Light}{LightSat} \cdot e^{-Extinction \cdot DepthTop}}$$
(34)

where:

Light = photosynthetically active radiation (ly/d); and LightSat = light saturation level for photosynthesis (ly/d).

Healthy blue-green algae tend to float. Therefore, if the nutrient limitation for blue-greens is greater than 0.25 (Equation (43)) and the wind is less than 3 m/s then *DepthBottom* for blue-greens is set to 0.25 m to account for buoyancy due to gas vacuoles. Otherwise it is set to 3 m to represent downward transport by Langmuir circulation. Other phytoplankton are considered to occupy all the well mixed layer. Under the ice, phytoplankton are represented as occurring in the top 2 m (cf. LeCren and Lowe-McConnell, 1980). As discussed in Section 3.5, light is decreased to 33% of incident radiation if ice cover is predicted.

Approximately half the incident solar radiation is photosynthetically active (Edmondson, 1956):

$$Light = Solar \cdot 0.5 \tag{35}$$

where:

Solar = average daily light intensity (ly/d), see (22).

The light-limitation function represents both limitation for suboptimal light intensity and photoinhibition at high light intensities (**Figure 25**). However, when the photoperiod for all but the highest latitudes is factored in, photoinhibition disappears (**Figure 26**). When considered over the course of the year, photoinhibition can occur in very clear, shallow systems during summer mid-day hours (**Figure 27**), but it usually is not a factor when considered over 24 hours (**Figure 28**).

The extinction coefficient for pure water varies considerably in the photosynthetically-active 400-700 nm range (Wetzel, 1975, p. 55); the value of 0.016 (1/m) is used, corresponding to the extinction of green light. In many models dissolved organic matter and suspended sediment are not considered separately, so a much larger extinction coefficient is used for "water" than in AQUATOX. The attenuation coefficients have units of 1/m-(g/m³) because they represent the amount of extinction caused by a given concentration (Table 6).

Figure 25
Instantaneous Light Response Function

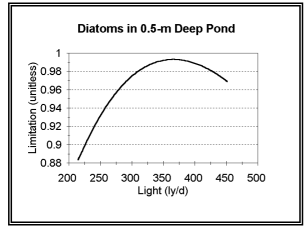


Figure 26Daily Light Response Function

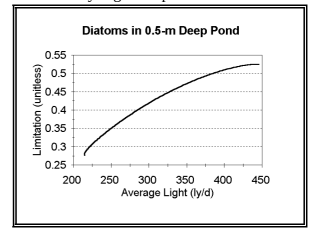


Figure 27Mid-day Light Limitation

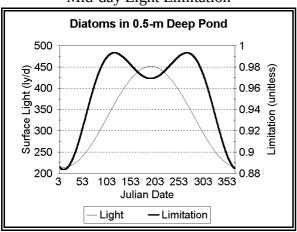


Figure 28Daily Light Limitation

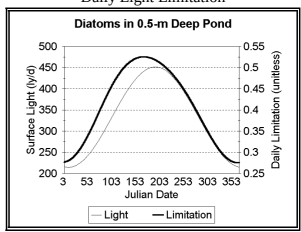


Table 6. Light Extinction and Attenuation Coefficients

WaterExtinction	0.016 1/m	Wetzel, 1975
ECoeffPhyto _{diatom}	$0.014 \text{ 1/m-(g/m}^3)$	Collins and Wlosinski, 1980
ECoeffPhyto _{blue-green}	$0.099 \text{ 1/m-(g/m}^3)$	Megard et al., 1979 (calc.)
ECoeffDOM	$0.03 \text{ 1/m-(g/m}^3)$	Effler et al., 1985 (calc.)
ECoeffPOM	$0.12 \text{ 1/m-(g/m}^3)$	Verduin, 1982
ECoeffSed	$0.03 \text{ 1/m-(g/m}^3)$	McIntire and Colby, 1978

The Secchi depth, the depth at which a Secchi disk disappears from view, is a commonly used indication of turbidity. It is computed as (Straškraba and Gnauck, 1985):

$$Secchi = \frac{1.9}{Extinction} \tag{36}$$

Secchi = Secchi depth (m).

This relationship could also be used to back-calculate an overall Extinction coefficient if only the Secchi depth is known for a site.

As a verification of the extinction computations, the calculated and observed Secchi depths were compared for Lake George, New York. The Secchi depth is estimated to be 8.3 m in Lake George, based on site data for the various components (**Figure 29**). This compares favorably with observed values of 7.5 to 11 (Clifford, 1982).

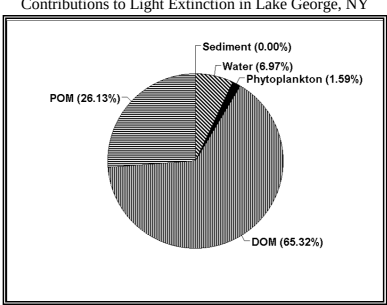


Figure 29
Contributions to Light Extinction in Lake George, NY

Nutrient Limitation

There are several ways that nutrient limitation has been represented in models. Algae are capable of taking up and storing sufficient nutrients to carry them through several generations, and models have been developed to represent this. However, if the timing of algal blooms is not critical, intracellular storage of nutrients can be ignored, constant stoichiometry can be assumed, and the model is much simpler. Therefore, based on the efficacy of this simplifying assumption, nutrient limitation by external nutrient concentrations is used in AQUATOX, as in many other models (for example, Chen, 1970; Parker, 1972; Lassen and Nielsen, 1972; Larsen et al., 1974; Park et al., 1974; Chen and Orlob, 1975; Patten et al., 1975; Environmental Laboratory, 1982; Ambrose et al., 1991).

For an individual nutrient, saturation kinetics is assumed, using the Michaelis-Menten or Monod equation (**Figure 30**); this approach is founded on numerous studies (cf. Hutchinson, 1967):

$$PLimit = \frac{Phosphorus}{Phosphorus + KP}$$
 (37)

$$NLimit = \frac{Nitrogen}{Nitrogen + KN}$$
 (38)

$$CLimit = \frac{Carbon}{Carbon + KCO2}$$
 (39)

where:

PLimit = limitation due to phosphorus (unitless); Phosphorus = available soluble phosphorus (gP/m³);

KP = half-saturation constant for phosphorus (gP/m^3);

NLimit = limitation due to nitrogen (unitless);
Nitrogen = available soluble nitrogen (gN/m³);

KN = half-saturation constant for nitrogen (gN/m³);
 CLimit = limitation due to inorganic carbon (unitless);
 Carbon = available dissolved inorganic carbon (gC/m³); and

KCO2 = half-saturation constant for carbon (gC/m³).

Nitrogen fixation in blue-green algae is handled by setting *NLimit* to 1.0 if *Nitrogen* is less than half the *KN* value. Otherwise, it is assumed that nitrogen fixation is not operable, and *NLimit* is computed as for the other algae.

Concentrations must be expressed in terms of the chemical element; therefore, the concentration of the compound is corrected for the molar weight of the element:

$$Phosphorus = P2PO4 \cdot Phosphate$$
 (40)

$$Carbon = C2CO2 \cdot CO2 \tag{41}$$

$$Nitrogen = N2NH4 \cdot Ammonia + N2NO3 \cdot Nitrate$$
 (42)

where:

P2PO4 = ratio of phosphorus to phosphate (0.33); Phosphate = available soluble phosphate (g/m³); N2NH4 = ratio of nitrogen to ammonia (0.78);

Ammonia = available ammonia (g/m^3) ;

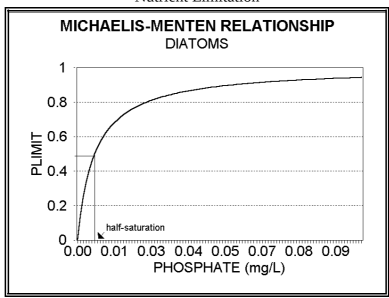
N2NO3 = ratio of nitrogen to nitrate (0.23);

C2CO2 = ratio of carbon to carbon dioxide (0.27); and

CO2 = inorganic carbon (g/m³).

All these conversions are built into AQUATOX.

Figure 30Nutrient Limitation



Like many models (for example, Larsen et al., 1973; Baca and Arnett, 1976; Scavia et al., 1976; Smith, 1978; Bierman et al., 1980; Park et al., 1980; Johanson et al., 1980; Grenney and Kraszewski, 1981; Ambrose et al., 1991), AQUATOX uses the minimum limiting nutrient, whereby the Monod equation is evaluated for each nutrient, and the factor for the nutrient that is most limiting at a particular time is used:

$$NutrLimit = min(PLimit, NLimit, CLimit)$$
 (43)

where:

NutrLimit = reduction due to limiting nutrient (unitless).

Alternative formulations used in other models include multiplicative and harmonic-mean constructs, but the minimum limiting nutrient construct is well-founded in laboratory studies with individual species.

Current Limitation

Because they are fixed in space, periphyton and macrophytes also are limited by slow currents that do not replenish nutrients and carry away senescent biomass. Based on the work of McIntire (1973) and Colby and McIntire (1978), a factor relating photosynthesis to current velocity is used for periphyton and macrophytes:

$$VLimit = \min(1, RedStillWater + \frac{VelCoeff \cdot Velocity}{1 + VelCoeff \cdot Velocity})$$
(44)

VLimit = limitation or enhancement due to current velocity (unitless); RedStillWater = reduction in photosynthesis in absence of current (unitless);

VelCoeff = empirical proportionality coefficient for velocity (0.057, unitless); and

Velocity = flow rate (m/s), see (169).

VLimit has a minimum value for photosynthesis in the absence of currents and increases asymptotically to a maximum value for optimal current velocity (**Figure 31**). In high currents entrainment can limit periphyton; see (60). The value of *RedStillWater* depends on the circumstances under which the maximum photosynthesis rate was measured; if *PMax* was measured in still water then *RedStillWater* = 1, otherwise a value of 0.2 is appropriate (Colby and McIntire, 1978).

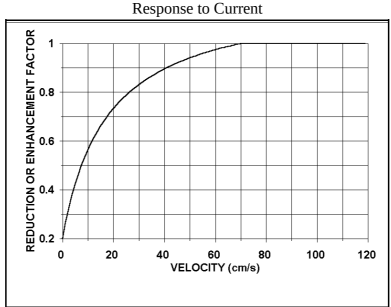


Figure 31
Response to Current

Adjustment for Suboptimal Temperature

AQUATOX uses a general but complex formulation to represent the effects of temperature. All organisms exhibit a nonlinear, adaptive response to temperature changes (the so-called Stroganov function). Process rates other than respiration increase as the ambient temperature increases until the optimal temperature for the organism is reached; beyond that optimum, process rates decrease until the lethal temperature is reached. This effect is represented by a complex algorithm developed by O'Neill et al. (1972) and modified slightly for application to aquatic systems (Park et al., 1974). An intermediate variable *VT* is computed first; it is the ratio of the difference between the maximum

temperature at which a process will occur and the ambient temperature over the difference between the maximum temperature and the optimal temperature for the process:

$$VT = \frac{(TMax + Acclimation) - Temperature}{(TMax + Acclimation) - (TOpt + Acclimation)}$$
(45)

where:

ambient water temperature (°C);

Temperature = TMax = maximum temperature at which process will occur (°C); optimal temperature for process to occur (°C); and *TOpt* Acclimation = temperature acclimation (°C), as described below.

Acclimation to changing temperature is accounted for with a modification developed by Kitchell et al. (1972):

Acclimation =
$$XM \cdot [1 - e^{(-KT \cdot ABS(Temperature - TRef))}]$$
 (46)

where:

XMmaximum acclimation allowed (°C);

coefficient for decreasing acclimation as temperature approaches $T_{\it ref}$ (unitless); KT

function to obtain absolute value; and ABS

TRef ="adaptation" temperature below which there is no acclimation (°C).

The mathematical sign of the variable Acclimation is negative if the ambient temperature is below the temperature at which there is no acclimation; otherwise, it is positive.

If the variable VT is less than zero, in other words, if the ambient temperature exceeds (TMax + Acclimation), then the suboptimal factor for temperature is set equal to zero and the process stops. Otherwise, the suboptimal factor for temperature is calculated as (Park et al., 1974):

$$TCorr = VT^{XT} \cdot e^{(XT \cdot (1-VT))} \tag{47}$$

where:

$$XT = \frac{WT^2 \cdot (1 + \sqrt{1 + 40/YT})^2}{400} \tag{48}$$

where:

$$WT = \ln(Q10) \cdot ((TMax + Acclimation) - (TOpt + Acclimation))$$
 (49)

and,

$$YT = \ln(Q10) \cdot ((TMax + Acclimation) - (TOpt + Acclimation) + 2)$$
 (50)

where:

Q10 = slope or rate of change per 10° C temperature change (unitless).

This well-founded, robust algorithm for *Tcorr* is used in AQUATOX to obtain reduction factors for suboptimal temperatures for all biologic processes in animals and plants, with the exception of algal respiration. By varying the parameters, organisms with both narrow and broad temperature tolerances can be represented (**Figure 32**, **Figure 33**).

Figure 32Temperature Response of Blue-Greens

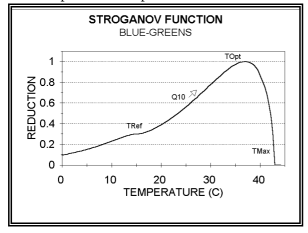
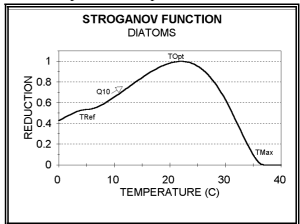


Figure 33Temperature Response of Diatoms



Algal Respiration

Endogenous or dark respiration is the metabolic process whereby oxygen is taken up by plants for the production of energy for maintenance and carbon dioxide is released (Collins and Wlosinski, 1983). Although it is normally a small loss rate for the organisms, it has been shown to be exponential with temperature (Aruga, 1965). Riley (1963, see also Groden, 1977) derived an equation representing this relationship. Based on data presented by Collins (1980), maximum respiration is constrained to 60% of photosynthesis. Laboratory experiments in support of the CLEANER model confirmed the empirical relationship and provided additional evidence of the correct parameter values (Collins, 1980), as demonstrated by **Figure 34**:

Respiration =
$$Resp0 \cdot e^{(TResp \cdot Temperature)} \cdot Biomass$$
 (51)

Respiration = dark respiration (g/m 3 ·d); Resp0 = respiration rate at 0°C (g/g·d);

TResp = exponential temperature coefficient (unitless);

Temperature = ambient water temperature (°C); and

Biomass = plant biomass (g/m³).

This construct also applies to macrophytes.

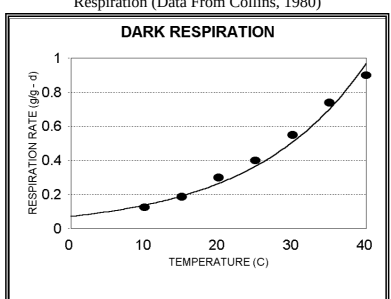


Figure 34Respiration (Data From Collins, 1980)

Photorespiration

Algal excretion, also referred to as photorespiration, is the release of photosynthate (dissolved organic material) and carbon dioxide that occurs in the presence of light. Environmental conditions that inhibit cell division but still allow photoassimilation result in release of organic compounds. This is especially true for both low and high levels of light (Fogg et al., 1965; Watt, 1966; Nalewajko, 1966; Collins, 1980). AQUATOX uses an equation modified from one by Desormeau (1978) that is the inverse of the light limitation:

$$Excretion = KResp \cdot LightStress \cdot Photosynthesis$$
 (52)

where:

Excretion = release of photosynthate $(g/m^3 \cdot d)$;

KResp = coefficient of proportionality between excretion and photosynthesis at

optimal light levels (unitless); and

Photosynthesis = photosynthesis ($g/m^3 \cdot d$), see (26),

and where:

$$LightStress = 1 - LtLimit$$
 (53)

where:

LtLimit= light limitation for a given plant (unitless), see (28).

It is a continuous function (**Figure 35**) and has a tendency to overestimate excretion slightly at light levels close to light saturation where experimental evidence suggests a constant relationship (Collins, 1980). The construct for photorespiration also applies to macrophytes.

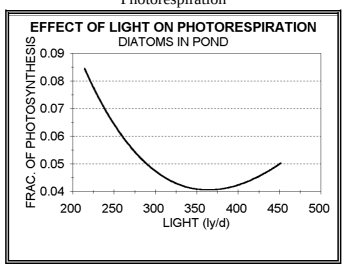


Figure 35 Photorespiration

Algal Mortality

Nonpredatory algal mortality can occur as a response to toxic chemicals (discussed in **Chapter 8**) and as a response to unfavorable environmental conditions. Phytoplankton under stress may suffer greatly increased mortality due to autolysis and parasitism (Harris, 1986). Therefore, most phytoplankton decay occurs in the water column rather than in the sediments (DePinto, 1979). The rapid remineralization of nutrients in the water column may result in a succession of blooms (Harris, 1986). Sudden changes in the abiotic environment may cause the algal population to crash; stressful changes include nutrient depletion, unfavorable temperature, and damage by light (LeCren and Lowe-McConnell, 1980). These are represented by a mortality term in AQUATOX that includes toxicity, high temperature (Scavia and Park, 1976) and combined nutrient and light limitation (Collins and Park, 1989):

$$Mortality = (KMort + ExcessT + Stress) \cdot Biomass + Poisoned$$
 (54)

Mortality = nonpredatory mortality $(g/m^3 \cdot d)$;

Poisoned = mortality rate due to toxicant $(g/g \cdot d)$, see ((269));

Kmort = intrinsic mortality rate due to high temperature $(g/g \cdot d)$; and

 $Biomass = plant biomass (g/m^3),$

and where:

$$ExcessT = \frac{e^{(Temperature - TMax)}}{2}$$
 (55)

and:

$$Stress = 1 - e^{-EMort \cdot (1 - (NutrLimit \cdot LtLimit))}$$
 (56)

where:

ExcessT = factor for high temperatures (g/g-d); TMax = maximum temperature tolerated ($^{\circ}$ C);

Stress = factor for suboptimal light and nutrients (g/g-d),

Emort = approximate maximum fraction killed per day; if total limitation then

value of 2 = doubled mortality (g/g-d);

NutrLimit = reduction due to limiting nutrient (unitless), see (43)

LtLimit = light limitation (unitless), see (29).

Exponential functions are used so that increasing stress leads to rapid increases in mortality, especially with high temperature where mortality is 50% per day at the *TMax* (**Figure 37**), and, to a much lesser degree, with suboptimal nutrients and light (**Figure 36**). This simulated process is responsible in part for maintaining realistically high levels of detritus in the simulated water body. Low temperatures are assumed not to affect algal mortality.

Figure 37Mortality Due To High Temperatures

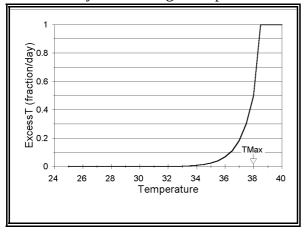
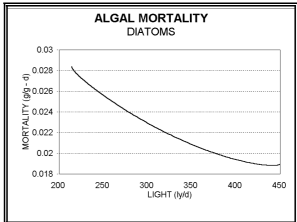


Figure 36Mortality Due To Light Limitation



Sinking

Sinking of phytoplankton, either between layers or to the bottom sediments, is modeled as a function of physiological state, similar to mortality. Phytoplankton that are not stressed are considered to sink at given rates, which are based on field observations and implicitly account for the effects of averaged water movements (cf. Scavia, 1980). Sinking also is represented as being impeded by turbulence associated with higher discharge:

$$Sink = \frac{KSed}{Depth} \cdot \frac{MeanDischarge}{Discharge + 0.001} \cdot SedAccel \cdot Biomass$$
 (57)

where:

Sink = phytoplankton loss due to settling (g/m³·d);

Ksed = intrinsic settling rate (m/d);

Depth = depth of water or, if stratified, thickness of layer (m);

MeanDischarge = mean annual discharge (m³/d);

Discharge = daily discharge (m^3/d), see Table 1; and

Biomass = phytoplankton biomass (g/m³).

As the phytoplankton are stressed by toxicants and suboptimal light, nutrients, and temperature, the model computes an exponential increase in sinking (**Figure 38**), as observed by Smayda (1974), and formulated by Collins and Park (1989):

$$SedAccel = e^{ESed \cdot (1 - LtLimit \cdot NutrLimit \cdot TCorr \cdot FracPhoto)}$$
(58)

where:

SedAccel = increase in sinking due to physiological stress (unitless);

ESed = exponential settling coefficient (unitless);

LtLimit = light limitation (unitless), see (28);

NutrLimit = nutrient limitation (unitless), see (43); and

FracPhoto = reduction factor for effect of toxicant on photosynthesis (unitless), see

(271);

TCorr = temperature limitation (unitless), see (47).

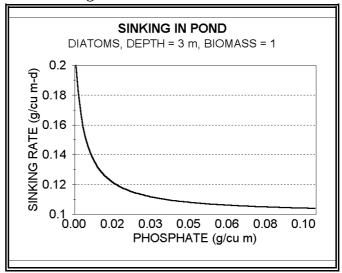


Figure 38Sinking as a Function of Nutrient Stress

This allows the model to mimic high sedimentation loss associated with the crashes of phytoplankton blooms, as discussed by Harris (1986). The equation is parameterized so that the sinking rate doubles as photosynthesis is totally limited, although that can be edited by the user.

Washout and Entrainment

Phytoplankton are subject to downstream drift. In streams and in lakes and reservoirs with low retention times this may be a significant factor in reducing or even precluding phytoplankton populations (LeCren and Lowe-McConnell, 1980). The process is modeled as a simple function of discharge:

$$Washout_{phytoplankton} = \frac{Discharge}{Volume} \cdot Biomass$$
 (59)

where:

Washout = loss due to downstream drift (g/m³·d), Discharge = daily discharge (m³/d), see Table 1; Volume = volume of site (m³), see (1) and Biomass = biomass of phytoplankton (g/m³).

Periphyton (and macrophytes, as discussed in the next section) also may be subject to entrainment and transport as they outgrow their substrate and as discharge increases (McIntire, 1968, 1973):

$$Washout_{periphyton, macrophytes} = Entrainment \cdot \frac{Discharge}{Volume} \cdot Biomass$$
 (60)

Entrainment is a function of carrying capacity; the formulation is based on McIntire (1973). As the biomass increases, additional biomass is entrained (**Figure 31**):

$$Entrainment = KCapLimit$$
 (61)

where:

Entrainment = fraction of biomass available for transport (unitless), and KCapLimit = limitation due to carrying capacity (unitless), see below.

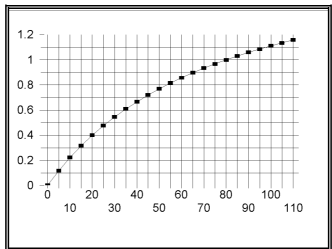
Because periphyton are limited by the area of substrate available, as the biomass approaches the carrying capacity of the substrate, increasing quantities are dislodged and available for transport (**Figure 39**):

$$KCapLimit = 1 - \frac{KCap - Biomass}{KCap}$$
 (62)

where:

KCap = carrying capacity of periphyton (g/m²).

Figure 39 Entrainment as a Function of Biomass



Chlorophyll a

Chlorophyll a is not simulated directly. However, because chlorophyll a is commonly measured in aquatic systems and because water quality managers are accustomed to thinking of it as an index of water quality, the model converts phytoplankton biomass estimates into approximate values for chlorophyll a. The ratio of carbon to chlorophyll a exhibits a wide range of values depending on the nutrient status of the algae (Harris, 1986); blue-green algae often have higher values (cf. Megard et al., 1979). AQUATOX uses a value of 45 μ gC/ μ g chlorophyll a for blue-greens and a value of 28 for other phytoplankton as reported in the documentation for WASP (Ambrose et al., 1991). The values are more representative for blooms than for static conditions, but managers are

usually most interested in the maxima. The results are presented as total chlorophyll a in μ g/L; therefore, the computation is:

$$Chlorophylla = \left(\frac{Biomass_{BlGr} \cdot CToOrg}{45} + \frac{\left(Biomass_{Diatom} + Biomass_{Oth}\right) \cdot CToOrg}{28}\right)$$

$$\cdot 1000$$
(63)

where:

 $Chlorophyll\ a =$ biomass as chlorophyll a ($\mu g/L$); Biomass biomass of given alga (mg/L);

= = CToOrg ratio of carbon to biomass (0.526, unitless); and

1000 conversion factor for mg to µg (unitless).

4.2 Macrophytes

Submersed aquatic vegetation or macrophytes can be an important component of shallow aquatic ecosystems. It is not unusual for the majority of the biomass in an ecosystem to be in the form of macrophytes during the growing season. Seasonal macrophyte growth, death, and decomposition can affect nutrient cycling, and detritus and oxygen concentrations. By forming dense cover, they can modify habitat and provide protection from predation for invertebrates and smaller fish (Howick et al., 1993); this function is represented in AQUATOX (see Figure 45). Macrophytes also provide direct and indirect food sources for many species of waterfowl, including swans, ducks, and coots (Jupp and Spence, 1977b).

AQUATOX represents macrophytes as occupying the littoral zone, that area of the bottom surface that occurs within the euphotic zone (see (11) for computation). Similar to periphyton, the compartment has units of g/m². In nature, macrophytes can be greatly reduced if phytoplankton blooms or higher levels of detritus increase the turbidity of the water (cf. Jupp and Spence, 1977a). Because the depth of the euphotic zone is computed as a function of the extinction coefficient (12), the area predicted to be occupied by macrophytes can increase or decrease depending on the clarity of the water.

The macrophyte equations are based on submodels developed for the International Biological Program (Titus et al., 1972; Park et al., 1974) and CLEANER models (Park et al., 1980) and for the Corps of Engineers' CE-QUAL-R1 model (Collins et al., 1985):

$$\frac{dBiomass}{dt} = Loading + Photosynthesis - Respiration - Excretion - Mortality - Predation - Washout$$
(64)

and:

$$Photosynthesis = PMax \cdot LtLimit \cdot VLimit \cdot TCorr \cdot Biomass \cdot FracLittoral \\ \cdot FracPhoto$$
(65)

dBiomass/dt = change in biomass with respect to time (g/m²·d);

Loading = loading of macrophyte, usually used as a "seed" $(g/m^2 \cdot d)$;

Photosynthesis = rate of photosynthesis (g/m 2 ·d); Respiration = respiratory loss (g/m 2 ·d), see (51);

Excretion = excretion or photorespiration(g/m²·d), see (52); Mortality = nonpredatory mortality (g/m²·d), see (66);

Predation = herbivory $(g/m^2 \cdot d)$, see (68);

Washout = loss due to entrainment $(g/m^2 \cdot d)$, see (60),

PMax = maximum photosynthetic rate (1/d), LtLimit = light limitation (unitless), see (29), VLimit = current limitation (unitless), see (44),

TCorr = correction for suboptimal temperature (unitless), see (47),

FracLittoral = fraction of bottom that is in the euphotic zone (unitless) see (11); and reduction factor for effect of toxicant on photosynthesis (unitless), see

(271).

They share many of the constructs with the algal submodel described above. Temperature limitation is modeled similarly, but with different parameter values. Light limitation also is handled similarly, using the Steele (1962) formulation; the application of this equation has been verified with laboratory data (Collins et al., 1985). Periphyton are epiphytic in the presence of macrophytes; by growing on the leaves they contribute to the light extinction for the macrophytes (Sand-Jensen, 1977). Extinction due to periphyton biomass is computed in AQUATOX, by inclusion in *LtLimit*. Nutrient limitation is not modeled at this time because macrophytes can obtain most of their nutrients from bottom sediments (Bristow and Whitcombe, 1971; Nichols and Keeney, 1976; Barko and Smart, 1980).

Simulation of respiration and excretion utilize the same equations as algae; excretion results in "nutrient pumping" because the nutrients are assumed to come from the sediments but are excreted to the water column. (Because nutrients are not explicitly modeled in bottom sediments, this can result in loss of mass balance, particularly in shallow ponds.) Non-predatory mortality is modeled similarly to algae as a function of suboptimal temperature and light. However, mortality is a function of low as well as high temperatures, and winter die-back is represented as a result of this control; the response is the inverse of the temperature limitation (**Figure 40**):

Mortality =
$$[Poisoned + (1 - e^{-EMort \cdot (1 - LtLimit \cdot TCorr)})] \cdot Biomass$$
 (66)

where:

Poisoned = mortality rate due to toxicant $(g/g \cdot d)$ (269), and

EMort = maximum mortality due to suboptimal conditions $(g/g \cdot d)$.

Sloughing of dead leaves can be a significant loss (LeCren and Lowe-McConnell, 1980); it is simulated as an implicit result of mortality (**Figure 41**).

Figure 40
Mortality as a Function of Temperature

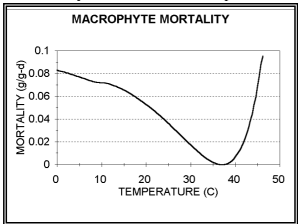
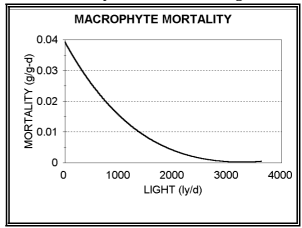


Figure 41Mortality as a Function of Light



Currents and wave agitation can both stimulate and retard macrophyte growth. These effects will be modeled in a future version. Similar to the effect on periphyton, water movement can stimulate photosynthesis in macrophytes (Westlake, 1967); the same function could be used for macrophytes as for periphyton, although with different parameter values. Jupp and Spence (1977b) have shown that wave agitation can severely limit macrophytes; time-varying entrainment eventually will be modeled when wave action is simulated.

4.3 Animals

Zooplankton, benthic invertebrates, benthic insects, and fish are modeled, with only slight differences in formulations, with a generalized animal submodel that is parameterized to represent different groups:

where:

dBiomass/dt = change in biomass of animal with respect to time (g/m³·d);

Load = biomass loading, usually from upstream $(g/m^3 \cdot d)$;

Consumption = consumption of food $(g/m^3 \cdot d)$, see (74);

Defecation = defecation of unassimilated food ($g/m^3 \cdot d$), see (73);

Respiration = respiration (g/m 3 ·d), see (76); Excretion = excretion (g/m 3 ·d), see (79);

Death = nonpredatory mortality (g/m³·d), see (80); Predation = predatory mortality (g/m³·d), see (75);

GameteLoss = loss of gametes during spawning (g/m³·d), see (84);

Washout = loss due to being carried downstream by washout and drift $(g/m^3 \cdot d)$,

see (87) and (88);

Migration = loss (or gain) due to vertical migration (g/m³·d), see (91);

Promotion = promotion to next size class or emergence $(g/m^3 \cdot d)$, see (92); and

Recruit = recruitment from previous size class $(g/m^3 \cdot d)$, see (92).

The change in biomass (**Figure 42**) is a function of a number of processes (**Figure 43**) that are subject to environmental factors, including biotic interactions. Similar to the way algae are treated, parameters for different species of invertebrates and fish are loaded and available for editing by means of the entry screens.

Consumption, Defecation, and Predation

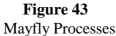
Several formulations have been used in various models to represent consumption of prey, reflecting the fact that there are different modes of feeding and that experimental evidence can be fit by any one of several equations (Mullin et al., 1975; Scavia, 1979; Straškraba and Gnauck, 1985).

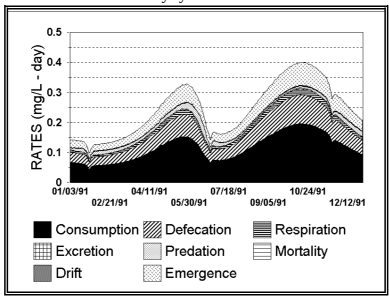
Ingestion is represented in AQUATOX by a maximum consumption rate, adjusted for ambient food and temperature conditions, and reduced for sublethal toxicant effects:

$$Ingestion_{prey, pred} = CMax_{pred} \cdot SatFeeding \cdot TCorr_{pred} \cdot ToxReduction \cdot Biomass_{pred}$$
 (68)

10 8 (1)6 (1)6 (2) 01/01/91 02/27/91 04/25/91 06/21/91 08/17/91 10/13/91 12/09/91 — Amphipods — Mayflies — Bass, YOY — Bass, adult — Minnows

Figure 42 Change in Animal Biomass in Stream





Many animals adjust their search or filtration in accordance with the concentration of prey; therefore, a saturation-kinetic term is used (Park et al., 1974, 1980; Scavia and Park, 1976):

$$SatFeeding = \frac{Preference_{prey, pred} \cdot Food}{\sum_{prey}(Preference_{prey, pred} \cdot Food) + FHalfSat_{pred}}$$
(69)

where:

 $Ingestion_{prey, pred} =$ ingestion of given prey by given predator (g/m³·d);

Biomass = concentration of organism $(g/m^3 \cdot d)$;

CMax = maximum feeding rate for predator (g/g·d);

TCorr = reduction factor for suboptimal temperature (unitless), see **Figure 32**;

Preference = preference of predator for prey (unitless);

Food = available food (g/m³);

FHalfSat = half-saturation constant for feeding by a predator (g/m³); and ToxReduction = reduction due to effects of toxicant (see Eq. (274), unitless).

The food actually available to a predator may be reduced in two ways:

$$Food = (Biomass_{prey} - BMin_{pred}) \cdot Refuge$$
 (70)

where:

BMin = minimum prey biomass needed to begin feeding (g/m^3) ; and *Refuge* = reduction factor for prey hiding in macrophytes (unitless).

Search or filtration may virtually cease below a minimum prey biomass (*BMin*) to conserve energy (**Figure 44**), so that a minimum food level is incorporated (Parsons et al., 1969; Steele, 1974; Park et al., 1974; Scavia and Park, 1976; Scavia et al., 1976; Steele and Mullin, 1977). However, cladocerans (for example, *Daphnia*) must constantly filter because the filtratory appendages also serve for respiration; therefore, in these animals there is no minimum feeding level.

Macrophytes can provide refuge from predation; this is represented by a factor related to the macrophyte biomass that is original with AQUATOX (**Figure 45**):

$$Refuge = 1 - \frac{Biomass_{Macro}}{Biomass_{Macro} + HalfSat}$$
 (71)

where:

HalfSat = half-saturation constant (20, g/m³), and

 $Biomass_{Macro}$ = biomass of macrophyte (g/m³).

Figure 44Saturation-kinetic Consumption

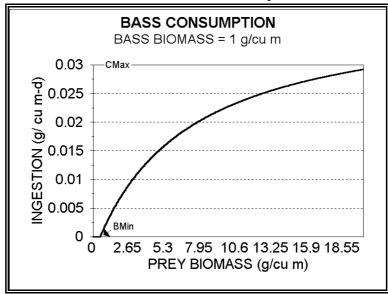
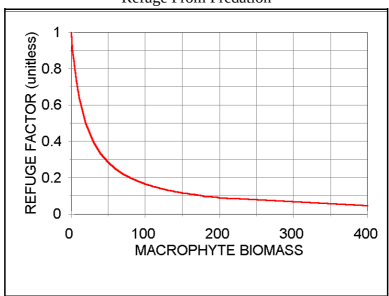


Figure 45Refuge From Predation



AQUATOX is a food-web model with multiple potential food sources. Passive size-selective filtering (Mullin, 1963; Lam and Frost, 1976) and active raptorial selection (Burns, 1969; Berman and Richman, 1974; Bogdan and McNaught, 1975; Brandl and Fernando, 1975) occur among aquatic organisms. Relative preferences are represented in AQUATOX by a matrix of preference parameters first proposed by O'Neill (1969) and used in several aquatic models (Bloomfield et al., 1973; Park et al., 1974; Canale et al., 1976; Scavia et al., 1976). Higher values indicate increased preference by a given predator for a particular prey compared to the preferences for all possible prey. In other words, the availability of the prey is weighted by the preference factor.

The preference factors are normalized so that if a potential food source is not modeled or is below the BMin value, the other preference factors are modified accordingly, representing adaptive preferences:

$$Preference_{prey,pred} = \frac{Pref_{prey,pred}}{SumPref}$$
 (72)

where:

 $Preference_{prey,pred}$ normalized preference of given predator for given prey

(unitless);

 $Pref_{prev, pred}$ initial preference value from the animal parameter screen

(unitless); and

SumPref sum of preference values for all food sources that are present

above the minimum biomass level for feeding during a

particular time step (unitless).

Similarly, different prey types have different potentials for assimilation by different predators. The fraction of ingested prey that is egested as feces or discarded (and which is treated as a source of detritus by the model, see (106)), is indicated by a matrix of egestion coefficients with the same structure as the preference matrix, so that defecation is computed as (Park et al., 1974):

$$Defecation_{pred} = \sum_{prey} (EgestCoeff_{prey, pred} + IncrEgest) \cdot Ingestion_{prey, pred})$$
 (73)

where:

total defecation for given predator (g/m³·d);

 $Detecation_{pred} = EgestCoeff_{prey, pred} = IncrEast$ fraction of ingested prey that is egested (unitless); and *IncrEgest* increased egestion due to toxicant (see Eq. (275), unitless).

Consumption of prey for a predator is also considered predation or grazing for the prey. Therefore, AQUATOX represents consumption as a source term for the predator and as a loss term for the prey:

$$Consumption_{pred} = \Sigma_{prey}(Ingestion_{prey, pred})$$
 (74)

$$Predation_{prey} = \Sigma_{pred}(Ingestion_{prey, pred})$$
 (75)

where

 $Consumption_{pred} = Predation_{prev} =$ total consumption rate by predator (g/m³·d); and

total predation on given prey (g/m³·d). Predation_{prev}

Respiration

Respiration can be considered as having two components (Park et al., 1979):

$$Respiration_{pred} = SpecDynAction_{pred} + Endogenous_{pred}$$
 (76)

where:

 $Respiration_{pred}$ = respiratory loss of predator (g/m³·d);

 $SpecDynAction_{pred}$ = respiratory loss due to activity (g/m³·d), see (78); and

Endogenous_{pred} = basal respiratory loss modified by temperature (g/m³·d); see

(77).

Basal or endogenous respiration is a rate at resting in which the organism is expending energy without uptake (as in overwintering), in contrast to the so-called specific dynamic action when the organism is moving, and consuming and digesting prey. AQUATOX simulates basal respiration as increasing with increasing temperature to a maximum value, using the adaptive temperature function (see Hewett and Johnson, 1992):

$$Endogenous_{pred} = EndogResp_{pred} \cdot TCorr_{pred} \cdot Biomass_{pred}$$
 (77)

where:

 $EndogResp_{pred}$ = basal respiration rate at 0° C for given predator (1/day); parameter

input by user as "Respiration Rate;"

 $TCorr_{pred}$ = Stroganov temperature function (unitless), see **Figure 32**; and

 $Biomass_{pred}$ = concentration of predator (g/m³).

As a simplification, specific dynamic action is represented as proportional to food assimilated

$$SpecDynAction_{pred} = KResp_{pred} \cdot (Consumption_{pred} - Defecation_{pred})$$
 (78)

(Hewett and Johnson, 1992; see also Kitchell et al., 1974; Park et al., 1974): where:

 $KResp_{pred}$ = proportion of assimilated energy lost to specific dynamic

action (unitless); parameter input by user as "Specific Dynamic

Action;"

Consumption_{pred} = ingestion (g/m 3 ·d); and

 $Defecation_{nred}$ = egestion of unassimilated food (g/m³·d).

Excretion

As respiration occurs, biomass is lost and nitrogen and phosphorus are excreted directly to the water (Horne and Goldman 1994); see (121) and (131). Ganf and Bla• ka (1974) have reported that this process is important to the dynamics of the Lake George, Uganda, ecosystem. Their data were

converted by Scavia and Park (1976) to obtain a proportionality constant relating excretion to respiration:

$$Excretion_{nred} = KExcr_{nred} \cdot Respiration_{nred}$$
 (79)

where:

excretion rate (g/m³·d);

 $Excretion_{pred} = KExcr_{pred} = Respiration_{pred} =$ proportionality constant for excretion:respiration (unitless); and

Respiration_{nred} respiration rate $(g/m^3 \cdot d)$.

Excretion is approximately 17 percent of respiration, which is not an important biomass loss term for animals, but it is important in nutrient recycling.

Nonpredatory Mortality

Nonpredatory mortality is a result of both environmental conditions and the toxicity of pollutants:

$$Death_{pred} = D_{pred} \cdot Biomass_{pred} + Poisoned_{pred}$$
 (80)

where:

 $Death_{pred} = D_{pred} =$ nonpredatory mortality (g/m³·d);

environmental mortality rate; the maximum value of three

computations, (81), (82), and (83), is used (1/d);

biomass of given animal (g/m³); and Biomass_{pred}

mortality due to toxic effects (g/m³·d), see (269). Poisoned

Under normal conditions a baseline mortality rate is used:

$$D_{pred} = KMort_{pred}$$
 (81)

where:

KMort_{pred} = normal nonpredatory mortality rate (1/d).

An exponential function is used for temperatures above the maximum (**Figure 46**):

$$D_{pred} = KMort_{pred} + \frac{e^{Temperature - TMax_{pred}}}{2}$$
 (82)

where:

ambient water temperature (°C); and Temperature = $TMax_{pred} =$ maximum temperature tolerated (°C).

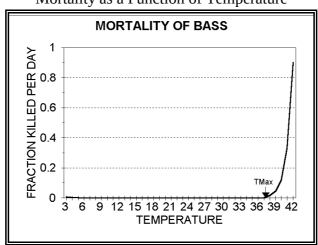


Figure 46Mortality as a Function of Temperature

The lower lethal temperature is often 0°C (Leidy and Jenkins, 1976), so it is ignored at this time. Total mortality is assumed when dissolved oxygen drops below 1 g/m³, recognizing that the predicted level is an average for the entire water column or epilimnetic or hypolimnetic segment:

$$Dead = 1.0 \text{ if } Oxygen < 1.0 \tag{83}$$

Gamete Loss and Recruitment

Eggs and sperm can be a significant fraction of adult biomass; in bluegills these can be 13 percent and 5 percent, respectively (Toetz, 1967), giving an average of 9 percent if the proportion of sexes is equal. Because only a small fraction of these gametes results in viable young when shed at the time of spawning, the remaining fraction is lost to detritus in the model. The construct is modified from a formulation by Kitchell et al. (1974). As a simplification, rather than requiring species-specific spawning temperatures, it assumes that spawning occurs when the temperature first enters the range from six tenths the optimum temperature to 1° less than the optimal temperature. This is based on a comparison of the optimal temperatures with the species-specific spawning temperatures reported by Kitchell et al. (1974). Depending on the range of temperatures, this simplifying assumption usually will result in one or two spawnings per year in a temperate ecosystem, which may or may not be realistic.

If
$$(0.6 \cdot TOpt) < Temperature < (TOpt - 1.0)$$
 then

$$GameteLoss = (GMort + IncrMort) \cdot FracAdults \cdot PctGamete \cdot Biomass$$

$$else \ GameteLoss = 0$$
(84)

where:

Temperature = ambient water temperature (°C);

TOpt = optimum temperature (°C); GameteLoss = loss rate for gametes (g/m³·d);

GMort = gamete mortality (1/d);

IncrMort = increased gamete and embryo mortality due to toxicant (see Eq.

(276),1/d);

Biomass = biomass of predator (g/m³);

PctGamete = fraction of adult predator biomass that is in gametes (unitless); and

FracAdults = fraction of biomass that is adult (unitless).

As the biomass of a population reaches its carrying capacity, reproduction is usually reduced due to stress; this results in a population that is primarily adults. Therefore, the proportion of adults and the fraction of biomass in gametes are assumed to be at a maximum when the biomass is at the carrying capacity (**Figure 47**):

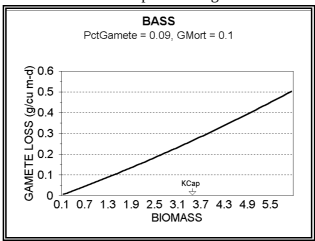
$$FracAdults = 1.0 - \left(\frac{Capacity}{KCap}\right)$$
 (85)

if Biomass > KCap then Capacity = 0 else Capacity = KCap - Biomass

where:

KCap = carrying capacity (g/m³).

Figure 47Correction for Population-Age Structure



Spawning in large gamefish results in an increase in the biomass of small gamefish if both small and large size classes are of the same species. Gametes are lost from the large gamefish, and the small gamefish gain the viable gametes through recruitment:

$$Recruit = (1 - (GMort + IncrMort)) \cdot FracAdults \cdot PctGamete \cdot Biomass$$
 (86)

Recruit = biomass gained from successful spawning (g/m³·d).

Washout and Drift

Downstream transport is an important loss term for invertebrates. Zooplankton are subject to transport downstream similar to phytoplankton:

$$Washout = \frac{Discharge}{Volume} \cdot Biomass$$
 (87)

where:

Washout = loss of zooplankton due to downstream transport $(g/m^3 \cdot d)$;

Discharge = discharge (m³/d), see Table 1; Volume = volume of site (m³), see (1); and Biomass = biomass of invertebrate (g/m³).

Likewise, zoobenthos exhibit drift, which is detachment followed by washout, and it is represented by a construct that is original with AQUATOX:

$$Drift = \frac{Discharge}{Volume} \cdot Dislodge \cdot Biomass$$
 (88)

where:

Drift = loss of zoobenthos due to downstream drift $(g/m^3 \cdot d)$; and

Dislodge = fraction of biomass subject to drift per day (unitless), see (89) and (90).

Nocturnal drift is a natural phenomenon:

$$Dislodge = NormalDrift$$
 (89)

where:

NormalDrift = fraction of biomass subject to normal drift per day (unitless).

However, drift is greatly increased when zoobenthos are subjected to stress by sublethal and lethal doses of toxic chemicals (Muirhead-Thomson, 1987), and that is represented by a saturation-kinetic formulation:

$$Dislodge = \frac{Toxicant_{Water}}{Toxicant_{Water} + EC50Growth}$$
(90)

where:

 $Toxicant_{Water}$ = concentration of toxicant in water (g/m³); and

EC50Growth = concentration at which half the population is affected (g/m³).

Vertical Migration

When presented with unfavorable conditions, most animals will attempt to migrate to an adjacent area with more favorable conditions. The current version of AQUATOX, following the example of CLEANER (Park et al., 1980), assumes that zooplankton and fish will exhibit avoidance behavior by migrating vertically from an anoxic hypolimnion to the epilimnion. The construct calculates the absolute mass of the given group of organisms in the hypolimnion, then divides by the volume of the epilimnion to obtain the biomass being added to the epilimnion:

If
$$VSeg = Hypo$$
 and $Anoxic$

$$Migration = \frac{HypVolume \cdot Biomass_{pred, hypo}}{EpiVolume}$$
(91)

where:

VSeg = vertical segment; Hypo = hypolimnion;

Anoxic = boolean variable for anoxic conditions;

Migration = rate of migration (g/m³·d);

HypVolume = volume of hypolimnion (m³), see **Figure 15**; volume of epilimnion (m³), see **Figure 15**; and biomass of given predator in hypolimnion (g/m³).

This does not include horizontal migration or avoidance of toxicants and stressful temperatures.

Promotion

Although AQUATOX is an ecosystem model, promotion to the next size class is important in representing the emergence of aquatic insects, and therefore loss of biomass from the system, and in predicting bioaccumulation of hydrophobic organic compounds in larger fish. The model assumes that promotion is determined by the rate of growth. Growth is considered to be the sum of consumption and the loss terms other than mortality and migration; a fraction of the growth goes into promotion to the next size class (cf. Park et al., 1980):

where:

Promotion = rate of promotion $(g/m^3 \cdot d)$;

KPro = fraction of growth that goes to promotion or emergence (0.5, unitless);

Consumption = rate of consumption (g/m 3 ·d), see (74); Defecation = rate of defecation (g/m 3 ·d), see (73); Respiration = rate of respiration (g/m 3 ·d), see (76); and Excretion = rate of excretion (g/m 3 ·d), see (79).

This is a simplification of a complex response that depends on the mean weight of the individuals. However, simulation of mean weight would require modeling both biomass and numbers of individuals (Park et al., 1979, 1980), and that is beyond the scope of this model at present.

Insect emergence can be an important factor in the dynamics of an aquatic ecosystem. Often there is synchrony in the emergence; in AQUATOX this is assumed to be cued to temperature, and is represented by:

If
$$Temperature > (0.8 \cdot TOpt)$$
 and $Temperature < (TOpt - 1.0)$ then
$$EmergeInsect = 2 \cdot Promotion$$
(93)

where:

EmergeInsect = insect emergence (mg/L·d);

Temperature = ambient water temperature (°C); and

TOpt = optimum temperature ($^{\circ}$ C);

5. REMINERALIZATION

5.1 Detritus

The term "detritus" is used to include all non-living organic material and associated decomposers (bacteria and fungi); as such, it includes both particulate and dissolved material in the sense of Wetzel (1975), but it also includes the microflora and is analogous to "biodetritus" of Odum and de la Cruz (1963). Detritus is now modeled as eight compartments: refractory (resistant) dissolved, suspended, sedimented, and buried detritus; and labile (readily decomposed) dissolved, suspended, sedimented, and buried detritus (**Figure 48**). This disaggregation is considered necessary to provide more realistic simulations of bioavailability of toxicants, with orders-of-magnitude differences in partitioning, and biological oxygen demand, which depends largely on the decomposition rates. Buried detritus is considered to be taken out of active participation in the functioning of the ecosystem. In general, dissolved organic material is about ten times that of suspended particulate matter in lakes and streams (Saunders, 1980), and refractory compounds usually predominate; however, the proportions are modeled dynamically.

≪<mark>fm</mark>. detr. Labile Refractory detr. fm. Dissolved Dissolved decomp. colonization detr. colonization Refractory Labile fm. detr ingestion fm Suspended ingestion Suspended decomp. sedimentation sedimentation scour scour detr. colonization Refractory Labile fm. detr ingestion fm. ingestion Sediments Sediments decomp. burial burial exposure exposure Refractory Labile **Buried** Buried + connection to nutrients * connection to detritivores

Figure 48
Detritus Compartments in AQUATOX

The concentrations of detritus in these eight compartments are the result of several competing processes:

$$\frac{dSuspRefrDetr}{dt} = Loading + DetrFm - Colonization - Washout - Sedimentation - Ingestion + Scour + TurbDiff$$
(94)

$$\frac{dSuspLabDetr}{dt} = Loading + DetrFm + Colonization - Decomposition - Washout - Sedimentation - Ingestion + Scour ± TurbDiff$$
(95)

$$\frac{dDissRefrDetr}{dt} = Loading + DetrFm - Colonization - Washout \pm TurbDiff$$
 (96)

$$\frac{dDissLabDetr}{dt} = Loading + DetrFm - Decomposition - Washout \pm TurbDiff$$
 (97)

$$\frac{dSedRefrDetr}{dt} = Loading + DetrFm + Sedimentation + Exposure - Colonization - Ingestion - Scour - Burial$$
(98)

$$\frac{dSedLabileDetr}{dt} = Loading + DetrFm + Sedimentation + Colonization - Ingestion - Decomposition - Scour + Exposure - Burial$$
(99)

$$\frac{dBuriedRefrDetr}{dt} = Sedimentation + Burial - Scour - Exposure$$
 (100)

$$\frac{dBuriedLabileDetr}{dt} = Sedimentation + Burial - Scour - Exposure$$
 (101)

dSuspRefrDetr/dt = change in concentration of suspended refractory detritus with respect to time (g/m³·d);

dSuspLabileDetr/dt = change in concentration of suspended labile detritus with respect to time (g/m³·d);

dDissRefrDetr/dt = change in concentration of dissolved refractory detritus with respect to time (g/m³·d);

dDissLabDetr/dt = change in concentration of dissolved labile detritus with

respect to time (g/m³·d);

dSedRefrDetr/dt = change in concentration of sedimented refractory detritus with

respect to time (g/m³·d);

dSedLabileDetr/dt = change in concentration of sedimented labile detritus with

respect to time $(g/m^3 \cdot d)$;

dBuriedRefrDetr/dt = change in concentration of buried refractory detritus with

respect to time (g/m³·d);

dBuriedLabileDetr/dt = change in concentration of buried labile detritus with respect

to time $(g/m^3 \cdot d)$;

Loading = loading of given detritus from nonpoint and point sources, or

from upstream (g/m³·d);

DetrFm = detrital formation (g/m 3 ·d);

Colonization = colonization of refractory detritus by decomposers (g/m³·d),

see (108);

Decomposition = loss due to microbial decomposition (g/m^3 ·d), see (112);

Sedimentation = transfer from suspended to sedimented by sinking (g/m³·d), see

(117);

Scour = resuspension from sedimented, and occasionally from buried,

to suspended (g/m³·d), see (162);

Exposure = transfer from buried to sedimented by scour of overlying

sediments (g/m³·d);

Burial = transfer from sedimented to buried due to deposition of

sediments (g/m³·d), see (165);

Washout = loss due to being carried downstream (g/m^3 ·d), see (13);

Ingestion = loss due to ingestion by detritivores and filter feeders $(g/m^3 \cdot d)$,

see (68); and

TurbDiff = transfer between epilimnion and hypolimnion due to turbulent

diffusion $(g/m^3 \cdot d)$, see (19) and (20).

As a simplification, refractory detritus is considered not to decompose directly, but rather to be converted to labile detritus through microbial colonization. Labile detritus is then available for both decomposition and ingestion by detritivores (organisms that feed on detritus). Because detritivores digest microbes and defecate the remaining organic material, detritus has to be conditioned through microbial colonization before it is suitable food. Therefore, the assimilation efficiency for refractory material is usually set to 0.0, and the assimilation efficiency for labile material is increased accordingly. Sedimentation and scour, or resuspension, are opposite processes. In shallow systems there may be no long-term sedimentation (Wetzel et al., 1972), while in deep systems there may be little resuspension. In this version sedimentation is a function of flow, ice cover and, in very shallow water, wind based on simplifying assumptions. Burial, scour and exposure are applicable only in streams where they are keyed to the behavior of clay and silt. Scour as an explicit function of wave and current action is not implemented.

Detrital Formation

Detritus is formed in several ways: through mortality, gamete loss, sinking of phytoplankton, excretion and defecation:

$$DetrFm_{SuspRefrDetr} = \sum_{biota} (Mort2_{detr, biota} \cdot Dead_{biota})$$
 (102)

$$DetrFm_{DissRefrDetr} = \Sigma_{biota}(Mort2_{detr,biota} \cdot Dead_{biota}) + \Sigma_{biota}(Excr2_{detr,biota} \cdot Excretion)$$
 (103)

$$DetrFm_{DissLabileDetr} = \Sigma_{biota}(Mort2_{detr,biota} \cdot Dead_{biota}) + \Sigma_{biota}(Excr2_{detr,biota} \cdot Excretion)$$
 (104)

$$DetrFm_{SuspLabileDetr} = \Sigma_{biota}(Mort2_{detr,biota} \cdot Dead_{biota}) + \Sigma_{animals} GameteLoss$$
 (105)

$$DetrFm_{SedLabileDetr} = \Sigma_{pred}(Def2_{detr,pred} \cdot Defecation_{pred}) + \Sigma_{compartment}(Sink_{compartment})$$
 (106)

$$DetrFm_{SedRefrDetr} = \Sigma_{pred}(Def2_{detr,pred} \cdot Defecation_{pred}) + \Sigma_{compartment}(Sink_{compartment})$$
 (107)

where:

DetrFm = formation of detritus (g/m 3 ·d);

 $Mort2_{detr. biota}$ = fraction of given dead organism that goes to given detritus (unitless);

 $Excr2_{detr. biota}$ = fraction of excretion that goes to given detritus (unitless);

 $Dead_{biota}$ = death rate for organism (g/m³·d), see (80);

Excretion = excretion rate for organism $(g/m^3 \cdot d)$, see (52) and (79) for plants and

animals, respectively;

GameteLoss = loss rate for gametes (g/m³·d), see (84);

 $Def2_{detr, biota}$ = fraction of defecation that goes to given detritus (unitless);

 $Defecation_{pred}$ = defecation rate for organism (g/m³·d), see (73); and

Sink = sinking rates for labile and refractory portions of phytoplankton

 $(g/m^3 \cdot d)$, see (57).

A fraction of mortality, including sloughing of leaves from macrophytes, is assumed to go to refractory detritus; a much larger fraction goes to labile detritus. Excreted material goes to both refractory and labile detritus, while gametes are considered to be labile. Half the defecated material is assumed to be labile because of the conditioning due to ingestion and subsequent inoculation with bacteria in the gut (LeCren and Lowe-McConnell, 1980); fecal pellets sink rapidly (Smayda, 1971),

so defecation is treated as if it were directly to sediments. Phytoplankton that sink to the bottom are considered to become detritus; most are consumed quickly by zoobenthos (LeCren and Lowe-McConnell, 1980) and are not available to be resuspended.

Colonization

Refractory detritus is converted to labile detritus through microbial colonization. When bacteria and fungi colonize dissolved refractory organic matter, they are in effect turning it into particulate matter. Detritus is usually refractory because it has a deficiency of nitrogen compared to microbial biomass. In order for microbes to colonize refractory detritus, they have to take up additional nitrogen from the water (Saunders et al., 1980). Thus, colonization is nitrogen-limited, as well as being limited by suboptimal temperature, pH, and dissolved oxygen:

$$Colonization = ColonizeMax \cdot DecTCorr \cdot NLimit \cdot pHCorr \\ \cdot DOCorrection \cdot RefrDetr$$
 (108)

where:

Colonization = rate of conversion of refractory to labile detritus (g/m³·d); maximum colonization rate under ideal conditions (g/g·d); ColonizeMax =Nlimit

limitation due to suboptimal nitrogen levels (unitless), see (110);

the effect of temperature (unitless), see (109); *DecTCorr*

limitation due to suboptimal pH level (unitless), see (115); pHCorr

limitation due to suboptimal oxygen level (unitless), see (113); and *DOCorrection* = concentration of refractory detritus in suspension, sedimented, or RefrDetr

dissolved (g/m^3) .

Because microbial colonization and decomposition involves microflora with a wide range of temperature tolerances, the effect of temperature is modeled in the traditional way (Thomann and Mueller, 1987), taking the rate at an observed temperature and correcting it for the ambient temperature up to a user-defined, high maximum temperature, at which point it drops to 0:

$$DecTCorr = Theta^{Temp - TObs}$$
 where
 $Theta = 1.047$ if $Temp \ge 19^{\circ}$ else
 $Theta = 1.185 - 0.00729 \cdot Temp$ (109)

The resulting curve has a shoulder similar to the Stroganov curve, but the effect increases up to the maximum rate (Figure 49).

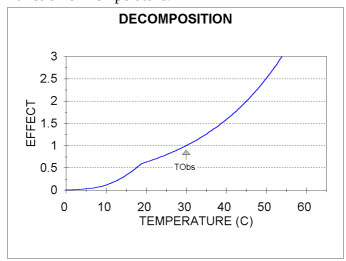


Figure 49. Colonization and Decomposition as a Function of Temperature.

The nitrogen limitation construct, which is original with AQUATOX, is computed by:

$$NLimit = \frac{N - MinN}{N - MinN + HalfSatN}$$
 (110)

$$N = N2NH4 \cdot Ammonia + N2NO3 \cdot Nitrate$$
 (111)

where:

N = total available nitrogen (g/m³);

MinN = minimum level of nitrogen for colonization (= 0.1 g/m³);

HalfSatN = half-saturation constant for nitrogen stimulation (= 0.15 g/m³);

N2NH4 = ratio of nitrogen to ammonia (= 0.78, unitless); and

N2NO3 = ratio of nitrogen to nitrate (= 0.23, unitless).

It is parameterized using an analysis of data presented by Egglishaw (1972) for Scottish streams. A maximum colonization rate of 0.007 (g/g·d) per day is used, based on McIntire and Colby (1978, after Sedell et al., 1975).

The rates of decomposition (or colonization) of refractory dissolved organic matter are comparable to those for particulate matter. Saunders (1980) reported values of 0.007 (g/g·d) for a eutrophic lake and 0.008 (g/g·d) for a tundra pond. Anaerobic rates were reported by Gunnison et al. (1985).

Decomposition

Decomposition is the process by which detritus is broken down by bacteria and fungi, yielding constituent nutrients, including nitrogen, phosphorus, and inorganic carbon. Therefore, it is a critical

process in modeling nutrient recycling. In AQUATOX, following a concept first advanced by Park et al. (1974), the process is modeled as a first-order equation with multiplicative limitations for suboptimal environmental conditions (see **section 4.1** for a discussion of similar construct for photosynthesis):

$$Decomposition = DecayMax \cdot DOCorrection \cdot DecTCorr \cdot pHCorr \cdot Detritus$$
 (112)

where:

Decomposition = loss due to microbial decomposition (g/m³·d);

DecayMax = maximum decomposition rate (g/g·d);

DOCorrection = correction for anaerobic conditions (unitless), see (113);

DecTCorr = the effect of temperature (unitless), see (109);

pHCorr = correction for suboptimal pH (unitless), see (115); and

Detritus = concentration of detritus, including dissolved but not buried (g/m^3) .

Note that biomass of bacteria is not explicitly modeled in AQUATOX. In some models (for example, EXAMS, Burns et al., 1982) decomposition is represented by a second-order equation using an empirical estimate of bacteria biomass. However, using bacterial biomass as a site constant constrains the model, potentially forcing the rate. Decomposers were modeled explicitly as a part of the CLEAN model (Clesceri et al., 1977). However, if conditions are favorable, decomposers can double in 20 minutes; this can result in stiff equations, adding significantly to the computational time. Ordinarily, decomposers will grow rapidly as long as conditions are favorable. The only time the biomass of decomposers might need to be considered explicitly is when a new organic chemical is introduced and the microbial assemblage requires time to become adapted to using it as a substrate.

The effect of temperature on biodegradation is represented by Equation (109), which also is used for colonization. The function for dissolved oxygen, formulated for AQUATOX, is:

$$DOCorrection = Factor + (1 - Factor) \cdot KAnaerobic$$
 (113)

where the predicted DO concentrations are entered into a Michaelis-Menten formulation to determine the extent to which degradation rates are affected by ambient DO concentrations (Clesceri, 1980; Park et al., 1982):

$$Factor = \frac{Oxygen}{HalfSatO + Oxygen}$$
 (114)

and:

Factor = Michaelis-Menten factor (unitless); KAnaerobic = decomposition rate at 0 g/m³ oxygen , Oxygen = dissolved oxygen concentration (g/m³); and HalfSatO = half-saturation constant for oxygen (g/m³). It accounts for both decreased (**Figure 50**) and increased (**Figure 51**) degradation rates under anaerobic conditions, with *KAnaerobic* having values less than one and greater than one, respectively. Detritus will always decompose more slowly under anaerobic conditions; but some organic chemicals, such as some halogenated compounds (Hill and McCarty, 1967), will degrade more rapidly. Half-saturation constants of 0.1 to 1.4 g/m³ have been reported (Bowie et al., 1985); a value of 0.5 g/m³ is used as a default.

Figure 50Correction for Dissolved Oxygen

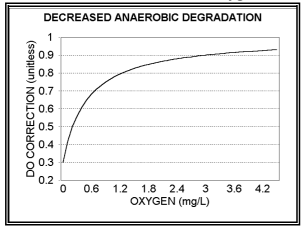
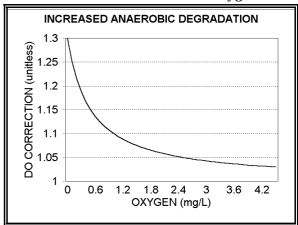


Figure 51Correction for Dissolved Oxygen



Another important environmental control on the rate of microbial degradation is pH. Most fungi grow optimally between pH 5 and 6 (Lyman et al., 1990), and most bacteria grow between pH 6 to about 9 (Alexander, 1977). Microbial oxidation is most rapid between pH 6 and 8 (Lyman et al., 1990). Within the pH range of 5 and 8.5, therefore, pH is assumed to not affect the rate of microbial degradation, and the suboptimal factor for pH is set to 1.0. In the absence of good data on the rates of biodegradation under extreme pH conditions, biodegradation is represented as decreasing exponentially beyond the optimal range (Park et al., 1980a; Park et al., 1982). If the pH is below the lower end of the optimal range, the following equation is used:

$$pHCorr = e^{(pH - pHMin)} (115)$$

where:

pH = ambient pH, and

pHMin = minimum pH below which limitation on biodegradation rate occurs.

If the pH is above the upper end of the optimal range for microbial degradation, the following equation is used:

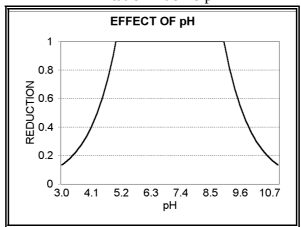
$$pHCorr = e^{(pHMmax - pH)} (116)$$

where:

pHMax = maximum pH above which limitation on biodegradation rate occurs.

These responses are shown in **Figure 52**.

Figure 52 Limitation Due To pH



Sedimentation

In this version, sedimentation of particulate detritus is modeled using simplifying assumptions. The constructs are intended to provide general responses to environmental factors, but they should not be considered as anything more than place holders for more realistic hydrodynamic functions to be incorporated in later versions.

$$Sedimentation = \frac{KSed}{Thick} \cdot Deaccel \cdot State$$
 (117)

where:

Sedimentation = transfer from suspended to sedimented by sinking $(g/m^3 \cdot d)$, see (117);

KSed = sedimentation rate (m/d);

Thick = depth of water or thickness of layer if stratified (m);

Deaccel = deceleration factor (unitless), see (118); and

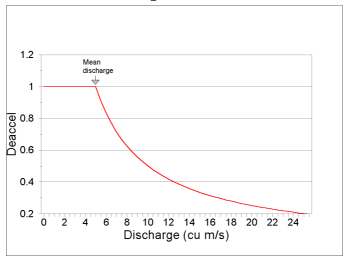
State = concentration of particulate detrital compartment (g/m^3) .

If the discharge exceeds the mean discharge then sedimentation is slowed proportionately (**Figure 53**):

If
$$TotDischarge > MeanDischarge$$
 then
$$Deaccel = \frac{MeanDischarge}{TotDischarge + 0.001}$$
else $Deaccel = 1.0$ (118)

TotDischarge = total epilimnetic and hypolimnetic discharge (m 3 /d); and MeanDischarge = mean discharge over the course of the simulation (m 3 /d).

Figure 53. Relationship of *Deaccel* to Discharge with a Mean Discharge of 5 m³/s.



If the depth of water is less than or equal to 1.0 m and wind speed is greater than or equal to 5.5 m/s then the sedimentation rate is negative, effectively becoming the rate of resuspension. If there is ice cover, then the sedimentation rate is doubled to represent the lack of turbulence.

5.2 Nitrogen

Two nitrogen compartments, ammonia and nitrate, are modeled (**Figure 54**). Nitrite occurs in very low concentrations and is rapidly transformed through nitrification and denitrification (Wetzel, 1975); therefore, it is modeled with nitrate. Likewise, un-ionized ammonia (NH₃) is not modeled as a separate state variable. Ammonia is assimilated by algae and macrophytes and is converted to nitrate as a result of nitrification:

$$\frac{dAmmonia}{dt} = Loading + Excrete + Decompose - Nitrify - Assim_{Ammonia} - Washout$$
 (119)

dAmmonia/dt = change in concentration of ammonia with time (g/m³·d);

Loading = loading of nutrient from inflow $(g/m^3 \cdot d)$;

Excrete = ammonia derived from excretion by animals $(g/m^3 \cdot d)$, see (121); Decompose = ammonia derived from decomposition of detritus $(g/m^3 \cdot d)$, see (120);

Nitrify = nitrification (g/m 3 ·d), see (127);

Assimilation = assimilation of nutrient by plants $(g/m^3 \cdot d)$, see (124) and (125); and loss of nutrient due to being carried downstream $(g/m^3 \cdot d)$, see (13).

Ammonia is a product of decomposition:

$$Decompose = \Sigma_{Detritus}(Org2Ammonia \cdot Decomposition_{Detritus})$$
 (120)

It is also excreted directly by organisms:

$$Excrete = \Sigma_{Biota}(Org2Ammonia \cdot Excretion_{Organism})$$
 (121)

where:

Org2Ammonia = ratio of ammonia to organic matter (unitless);

Decomposition = decomposition rate of given type of detritus, $(g/m^3 \cdot d)$, see

(112); and

Excretion = excretion rate of given organism $(g/m^3 \cdot d)$, see (79).

Nitrate is assimilated by plants and is converted to free nitrogen (and lost) through denitrification:

$$\frac{dNitrate}{dt} = Loading + AtmosDep + Nitrify - Denitrify - Assim_{Nitrate} - Washout$$
 (122)

where:

dNitrate/dt = change in concentration of nitrate with time (g/m³·d);

AtmosDep = atmospheric deposition (g/m 3 ·d); and

Denitrify = denitrification $(g/m^3 \cdot d)$.

Deposition directly from the atmosphere is:

$$AtmosDep = NAtmos \cdot \frac{Area}{Volume}$$
 (123)

where:

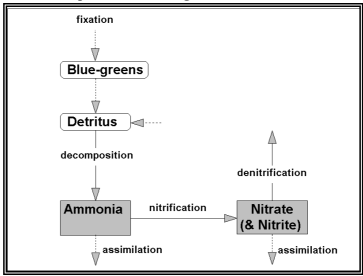
NAtmos = average observed atmospheric deposition rate (g/m²·d);

 $Area = area of site (m^2); and$

Volume = volume of water at site (m³).

Free nitrogen can be fixed by blue-green algae. Both nitrogen fixation and denitrification are subject to environmental controls and are difficult to model with any accuracy; therefore, the nitrogen cycle is represented with considerable uncertainty.

Figure 54Components of Nitrogen Remineralization



Assimilation

Nitrogen compounds are assimilated by plants as a function of photosynthesis in the respective groups (Ambrose et al., 1991):

$$Assimilation_{Ammonia} = \sum_{Plant} (Photosynthesis_{Plant} \cdot Uptake_{Nitrogen} \cdot NH4Pref)$$
 (124)

$$Assimilation_{Nitrate} = \sum_{Plant} (Photosynthesis_{Plant} \cdot Uptake_{Nitrogen} \cdot (1 - NH4Pref))$$
 (125)

where:

Assimilation = assimilation rate for given nutrient (g/m³·d); Photosynthesis = rate of photosynthesis (g/m³·d), see (26);

Uptake = fraction of photosynthate that is nutrient (unitless);

NH4Pref = ammonia preference factor (unitless),

Only 23 percent of nitrate is nitrogen, but 78 percent of ammonia is nitrogen. This results in an apparent preference for ammonia. The preference factor is calculated with an equation developed by Thomann and Fitzpatrick (1982) and cited and used in WASP (Ambrose et al., 1991):

$$NH4Pref = \frac{N2NH4 \cdot Ammonia \cdot N2NO3 \cdot Nitrate}{(KN + N2NH4 \cdot Ammonia) \cdot (KN + N2NO3 \cdot Nitrate)} + \frac{N2NH4 \cdot Ammonia \cdot KN}{(N2NH4 \cdot Ammonia + N2NO3 \cdot Nitrate) \cdot (KN + N2NO3 \cdot Nitrate)}$$

$$(126)$$

N2NH4 = ratio of nitrogen to ammonia (0.78); N2NO3 = ratio of nitrogen to nitrate (0.23);

KN = half-saturation constant for nitrogen uptake (g N/m³);

Ammonia = concentration of ammonia (g/m^3) ; and

Nitrate = concentration of nitrate (g/m^3) .

For algae other than blue-greens, *Uptake* is the Redfield (1958) ratio; although other ratios (cf. Harris, 1986) may be used by editing the parameter screen. At this time nitrogen-fixation by blue-greens is represented by using a smaller uptake ratio, thus "creating" nitrogen.

Nitrification and Denitrification

Nitrification is the conversion of ammonia to nitrite and then to nitrate by nitrifying bacteria. The maximum rate of nitrification is reduced by limitation factors for suboptimal dissolved oxygen and pH, similar to the way that decomposition is modeled, but using the more restrictive correction for suboptimal temperature used for plants and animals:

$$Nitrify = KNitri \cdot DOCorrection \cdot TCorr \cdot pHCorr \cdot Ammonia$$
 (127)

where:

Nitrify = nitrification rate $(g/m^3 \cdot d)$;

KNitri = maximum rate of nitrification (g nitrate/g ammonia);
 DOCorrection = correction for anaerobic conditions (unitless) see (113);
 TCorr = correction for suboptimal temperature (unitless); see (47);

pHCorr = correction for suboptimal pH (unitless), see (115);

Ammonia = concentration of ammonia (g/m^3) ; and

The nitrifying bacteria have narrow environmental optima; according to Bowie et al. (1985) they require aerobic conditions with a pH between 7 and 9.8, an optimal temperature of 30°, and minimum and maximum temperatures of 10° and 60° respectively (**Figure 55**, **Figure 56**).

Figure 55Response to pH, Nitrification

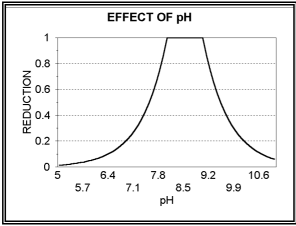
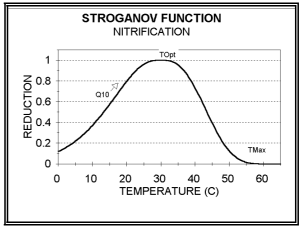


Figure 56Response to Temperature, Nitrification



In contrast, denitrification (the conversion of nitrate and nitrite to free nitrogen) is an anaerobic process, so that *DOCorrection* enhances the process (Ambrose et al., 1991):

$$Denitrify = KDenitri \cdot (1 - DOCorrection) \cdot TCorr \cdot pHCorr \cdot Nitrate$$
 (128)

where:

Denitrify = denitrification rate $(g/m^3 \cdot d)$;

KDenitri = maximum rate of denitrification (g ammonia/g nitrate); and

Nitrate = concentration of nitrate (g/m^3) .

Furthermore, it is accomplished by a large number of reducing bacteria under anaerobic conditions and with broad environmental tolerances (Bowie et al., 1985; **Figure 57**, **Figure 58**).

Figure 57 Response to pH, Denitrification

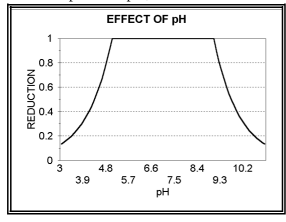
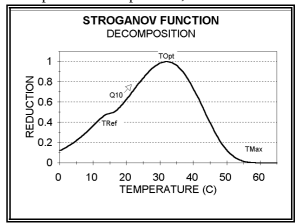


Figure 58Response to Temperature, Denitrification



5.3 Phosphorus

$$\frac{dPhosphate}{dt} = Loading \cdot FracAvail + AtmosDep + Excrete + Decompose - Assim_{Phosphate} - Washout$$
 (129)

The phosphorus cycle is much simpler than the nitrogen cycle. Decomposition, excretion, and assimilation are important processes that are similar to those described above:

$$AtmosDep = PAtmos \cdot \frac{Area}{Volume}$$
 (130)

$$Excrete = \Sigma_{Biota}(Org2Phosphate \cdot Excretion_{Biota})$$
 (131)

$$Decompose = \Sigma_{Detritus}(Org2Phosphate \cdot Decomposition_{Detritus})$$
 (132)

Assimilation =
$$\Sigma_{Plant}$$
 (Photosynthesis_{Plant} · Uptake_{Phosphorus}) (133)

where:

dPhosphate/dt = change in concentration of phosphate with time (g/m³·d);

Loading = loading of nutrient from inflow $(g/m^3 \cdot d)$;

FracAvail = fraction of phosphate loading that is available (unitless); AtmosDep = loading of nutrient directly from atmosphere ($g/m^3 \cdot d$); Excrete = phosphate derived from excretion by biota ($g/m^3 \cdot d$);

Decompose = phosphate derived from decomposition of detritus $(g/m^3 \cdot d)$;

Assimilation = assimilation by plants $(g/m^3 \cdot d)$;

Washout = loss due to being carried downstream (g/m 3 ·d), see (13); Patmos = average observed atmospheric deposition rate (g/m 2 ·d);

 $Area = area of site (m^2);$

Volume = volume of water at site (m³);

Org2Phosphate = ratio of phosphate to organic matter (unitless); Excretion = excretion rate for given organism (g/m 3 ·d), see (79);

Decomposition = decomposition rate for given detrital compartment (g/m 3 ·d), see (112);

Photosynthesis = rate of photosynthesis ($g/m^3 \cdot d$), see (26), and

Uptake = fraction of photosynthate that is phosphate (unitless).

At this time AQUATOX models only phosphate available for plants; a correction factor in the loading screen allows the user to scale total phosphate loadings to available phosphate. A future enhancement could be to consider phosphate precipitated with calcium carbonate, which would better represent the dynamics of marl lakes; however, that process is ignored in the current version. A default value is provided for average atmospheric deposition, but this should be adjusted for site conditions. In particular, entrainment of dust from tilled fields and new highway construction can cause significant increases in phosphate loadings. As with nitrogen, the uptake parameter is the Redfield (1958) ratio; it may be edited if a different ratio is desired (cf. Harris, 1986).

5.4 Dissolved Oxygen

Oxygen is an important regulatory endpoint; very low levels can result in mass mortality for fish and other organisms, mobilization of nutrients and metals, and decreased degradation of toxic organic materials. Dissolved oxygen is a function of reaeration, photosynthesis, respiration, decomposition, and nitrification:

$$\frac{dOxygen}{dt} = Loading + Reaeration + Photosynthesized - BOD - NitroDemand - Washout$$
 (134)

Photosynthesized =
$$O2Biomass \cdot \Sigma_{Plant}(Photosynthesis_{Plant})$$
 (135)

$$BOD = O2Biomass \cdot (\Sigma_{Detritus}(Decomposition_{Detritus}) + \Sigma_{Organisms}(Respiration_{Organisms}))$$
 (136)

$$NitroDemand = O2N \cdot Nitrify$$
 (137)

where:

dOxygen/dt = change in concentration of dissolved oxygen (g/m³·d);

Loading = loading from inflow $(g/m^3 \cdot d)$;

Reaeration = atmospheric exchange of oxygen (g/m³·d); Photosynthesized = oxygen produced by photosynthesis (g/m³·d); BOD = instantaneous biological oxygen demand (g/m³·d);

NitroDemand = oxygen taken up by nitrification $(g/m^3 \cdot d)$;

Washout = loss due to being carried downstream (g/m^3 ·d), see (13);

O2Biomass=ratio of oxygen to organic matter (unitless);Photosynthesis=rate of photosynthesis (g/m³·d), see (26), (65);Decomposition=rate of decomposition (g/m³·d), see (112);Respiration=rate of respiration (g/m³·d), see (76);O2N=ratio of oxygen to nitrogen (unitless); and

Nitrify = rate of nitrification (g $N/m^3 \cdot d$).

Reaeration is a function of the depth-averaged mass transfer coefficient *KReaer*, corrected for ambient temperature, multiplied by the difference between the dissolved oxygen level and the saturation level (cf. Bowie et al., 1985):

$$Reaeration = KReaer \cdot (O2Sat - Oxygen)$$
 (138)

where:

Reaeration = mass transfer of oxygen $(g/m^3 \cdot d)$;

KReaer = depth-averaged reaeration coefficient (1/d);

O2Sat = saturation concentration of oxygen (g/m³), see (147); and

Oxygen = concentration of oxygen (g/m^3) .

In standing water *KReaer* is computed as a minimum transfer velocity plus the effect of wind on the transfer velocity (Schwarzenbach et al., 1993) divided by the thickness of the mixed layer to obtain a depth-averaged coefficient (**Figure 59**):

$$KReaer = \frac{0.346 + 0.0346 \cdot Wind^2}{Thick}$$
 (139)

where:

Wind = wind velocity 10 m above the water (m/sec); and

Thick = thickness of mixed layer (m).

In streams, reaeration is a function of current velocity and water depth (**Figure 60**) following the approach of Covar (1978, see Bowie et al., 1985) and used in WASP (Ambrose et al., 1991). The decision rules for which equation to use are taken from the WASP5 code (Ambrose et al., 1991).

If *Vel* < 0.518:

$$TransitionDepth = 0$$
 (140)

else:

$$TransitionDepth = 4.411 \cdot Vel^{2.9135} \tag{141}$$

where:

Vel = velocity of stream (m/sec); and TransitionDepth = intermediate variable (m).

If *Depth* < 0.61 m, the equation of Owens et al. (1964, cited in Ambrose et al., 1991) is used:

$$KReaer = 5.349 \cdot Vel^{0.67} \cdot Depth^{-1.85}$$
 (142)

Depth = mean depth of stream (m).

Otherwise, if *Depth* is > *TransitionDepth*, the equation of O'Connor and Dobbins (1958, cited in Ambrose et al., 1991) is used:

$$KReaer = 3.93 \cdot Vel^{0.50} \cdot Depth^{-1.50}$$
 (143)

Else, if $Depth \leq TransitionDepth$, the equation of Churchill et al. (1962, cited in Ambrose et al., 1991) is used:

$$KReaer = 5.049 \cdot Vel^{0.97} \cdot Depth^{-1.67}$$
 (144)

In extremely shallow streams, especially experimental streams where depth is < 0.06 m, an equation developed by Krenkel and Orlob (1962, cited in Bowie et al. 1985) from flume data is used:

$$KReaer = \frac{234 \cdot (U \cdot Slope)^{0.408}}{H^{0.66}}$$

where:

U = velocity (fps);

Slope = longitudinal channel slope (m/m); and

H =water depth (ft).

If reaeration due to wind exceeds that due to current velocity, the equation for standing water is used. Reaeration is set to 0 if ice cover is expected (i.e., when the depth-averaged temperature < 3°C).

Figure 59Reaeration as a Function of Wind

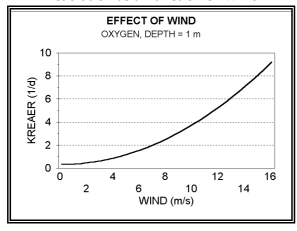
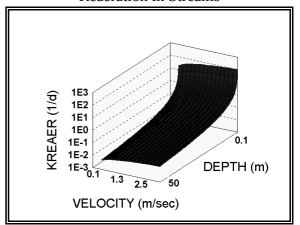


Figure 60
Reaeration in Streams



Reaeration is assumed to be representative of 20°C, so it is adjusted for ambient water temperature using (Thomann and Mueller 1987):

$$KReaer_T = KReaer_{20} \cdot Theta^{(Temperature - 20)}$$
 (146)

where:

 $KReaer_T$ = Reaeration coefficient at ambient temperature (1/d);

 $Kreaer_{20}$ = Reaeration coefficient for 20°C (1/d); Theta = temperature coefficient (1.024); and Temperature = ambient water temperature (°C).

Oxygen saturation, as a function of both temperature (**Figure 61**) and salinity (**Figure 62**), is based on Weiss (1970, cited in Bowie et al., 1985):

$$O2Sat = 1.4277 \cdot \exp[-173.4927 + \frac{24963.39}{TKelvin} + 143.3483 \ln\left(\frac{TKelvin}{100}\right) - 0.21849$$

$$\cdot TKelvin + S \cdot (-0.033096 + 0.00014259 \cdot TKelvin - 1.7 \cdot 10^{-7} \cdot \sqrt{TKelvin})]$$
(147)

where:

TKelvin = Kelvin temperature, and

S = salinity (ppt).

According to Bowie et al. (1985), it gives results that are not significantly different from those computed by the more complex APHA (1985) equations that are used in WASP (et al., 1993). At the present time salinity is set to 0; although, it has little effect on reaeration.

Figure 61
Saturation as a Function of Temperature

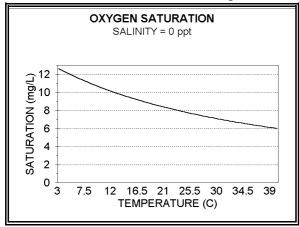
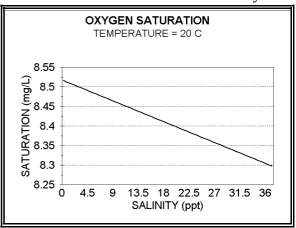


Figure 62
Saturation as a Function of Salinity



5.5 Inorganic Carbon

Many models ignore carbon dioxide as an ecosystem component (Bowie et al., 1985). However, it can be an important limiting nutrient. Similar to other nutrients, it is produced by decomposition and is assimilated by plants; it also is respired by organisms:

$$\frac{dCO2}{dt} = Loading + Respired + Decompose$$

$$- Assimilation - Washout \pm CO2AtmosExch$$
(148)

where:

Respired =
$$CO2Biomass \cdot \Sigma_{Organism}(Respiration_{Organism})$$
 (149)

Assimilation =
$$\Sigma_{Plant}(Photosynthesis_{Plant} \cdot UptakeCO2)$$
 (150)

$$Decompose = CO2Biomass \cdot \Sigma_{Detritus}(Decomp_{Detritus})$$
 (151)

and where:

dCO2/dt = change in concentration of carbon dioxide (g/m³·d); Loading = loading of carbon dioxide from inflow (g/m³·d); Respired = carbon dioxide produced by respiration (g/m³·d);
Decompose = carbon dioxide derived from decomposition (g/m³·d);
Assimilation = assimilation of carbon dioxide by plants (g/m³·d);
Washout = loss due to being carried downstream (g/m³·d), see (13);

CO2AtmosExch = interchange of carbon dioxide with atmosphere (g/m³·d);

co2Biomass = ratio of carbon dioxide to organic matter (unitless);

Respiration = rate of respiration (g/m³·d), see (76);
Decomposition = rate of decomposition (g/m³·d), see (112);
Photosynthesis = rate of photosynthesis (g/m³·d), see (26); and
UptakeCO2 = ratio of carbon dioxide to photosynthate (= 0.53).

Carbon dioxide also is exchanged with the atmosphere; this process is important, but is not instantaneous: significant undersaturation and oversaturation are possible (Stumm and Morgan, 1996). The treatment of atmospheric exchange is similar to that for oxygen:

$$CO2AtmosExch = KLiqCO2 \cdot (CO2Sat - CO2)$$
 (152)

In fact, the mass transfer coefficient is based on the well-established reaeration coefficient for oxygen, corrected for the difference in diffusivity of carbon dioxide as recommended by Schwarzenbach et al. (1993):

$$KLiqCO2 = KReaer \cdot \left(\frac{MolWtO2}{MolWtCO2}\right)^{0.25}$$
(153)

where:

CO2AtmosExch = interchange of carbon dioxide with atmosphere (g/m³·d); KLiqCO2 = depth-averaged liquid-phase mass transfer coefficient (1/d);

CO2 = concentration of carbon dioxide (g/m³);

CO2Sat = saturation concentration of carbon dioxide (g/m³), see (154);

KReaer = depth-averaged reaeration coefficient for oxygen (1/d), see (139)-

(146);

MolWtO2 = molecular weight of oxygen (=32); and MolWtCO2 = molecular weight of carbon dioxide (= 44).

Keying the mass-transfer coefficient for carbon dioxide to the reaeration coefficient for oxygen is very powerful in that the effects of wind (**Figure 63**) and the velocity and depth of streams can be represented, using the oxygen equations (Equations (139)- (144)).

EFFECT OF WIND

CARBON DIOXIDE, DEPTH = 1 m

10

8

P) 6

00

4

2

0

2

4

6

8

10

11

12

14

16

WIND (m/s)

Figure 63Carbon Dioxide Mass Transfer

Based on this approach, the predicted mass transfer under still conditions is 0.92, compared to the observed value of 0.89 ± 0.03 (Lyman et al., 1982). This same approach is used, with minor modifications, to predict the volatilization of other chemicals (see Section 7.5). Computation of saturation of carbon dioxide is based on the method in Bowie et al. (1985; see also Chapra and Reckhow, 1983) using Henry's law constant, with its temperature dependency (**Figure 64**), and the partial pressure of carbon dioxide:

$$CO2Sat = CO2Henry \cdot pCO2 \tag{154}$$

where:

$$CO2Henry = MCo2 \cdot 10^{\frac{2385.73}{TKelvin}} - 14.0184 + 0.0152642 \cdot TKelvin$$
 (155)

$$TKelvin = 273.15 + Temperature$$
 (156)

and where:

CO2Sat = saturation concentration of carbon dioxide (g/m³); CO2Henry = Henry's law constant for carbon dioxide (g/m³-atm)

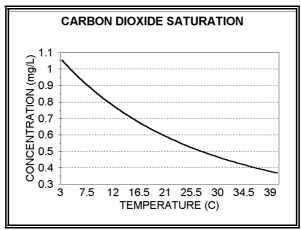
pCO2 = atmospheric partial pressure of carbon dioxide (= 0.00035);

MCO2 = mg carbon dioxide per mole (= 44000);

Tkelvin = temperature in °K, and

Temperature = ambient water temperature (°C).

Figure 64Saturation of Carbon Dioxide



6. INORGANIC SEDIMENTS¹

The sediment transport component of AQUATOX simulates scour, deposition and transport of sediments and calculates the concentration of sediments in the water column and sediment bed within a river reach. For running waters, the sediment is divided into three categories according to the particle size: 1) sand, with particle sizes between 0.062 to 2.0 millimeters (mm), 2) silt (0.004 - 0.062 mm), and 3) clay (0.00024 0.004 mm). Wash load (primarily clay and silt) is deposited or eroded within the channel reach depending on the daily flow regime. Sand transport is also computed within the channel reach. At present, inorganic sediments in standing water are computed based on total suspended solids loadings, described in section 6.3.

The river reach is assumed to be short and well mixed so that concentration does not vary longitudinally. Flow routing is not performed within the river reach. The daily average flow regime determines the amount of scour, deposition and transport of sediment. Scour, deposition and transport quantities are also limited by the amount of solids available in the bed sediments and the water column.

Inorganic sediments are important to the functioning of natural and perturbed ecosystems for several reasons. When suspended, they increase light extinction and decrease photosynthesis. When sedimented, they can temporarily or permanently remove toxicants from the active ecosystem through deep burial. Rapid sedimentation can adversely affect periphyton and some zoobenthos. Scour can also adversely affect periphyton and zoobenthos. The ratio of inorganic to organic sediments can be used as an indicator of aerobic or anaerobic conditions in the bottom sediments.

The mass of sediment in each of the three sediment size classes is a function of the previous mass, and the mass of sediment in the overlying water column lost through deposition, and gained through scour:

$$MassBed_{Sed} = MassBed_{Sed, t=-1} + (Deposit_{Sed} - Scour_{Sed}) \cdot Volume_{Water}$$
 (157)

where:

 $MassBed_{Sed}$ = mass of sediment in channel bed (kg);

 $MassBed_{Sed, t=-1}$ = mass of sediment in channel bed on previous day (kg); $Deposit_{Sed}$ = amount of suspended sediment deposited (kg/m³); $Scour_{Sod}$ = amount of silt or clay resuspended (kg/m³); and

 $Volume_{Water}$ = volume of stream reach (m³).

The volumes of the respective sediment size classes are calculated as:

¹ Original riverine version contributed by Rodolfo Camacho of Abt Associates Inc.; not validated

$$Volume_{Sed} = \frac{MassBed_{Sed}}{Rho_{Sed}}$$
 (158)

 $Volume_{Sed}$ volume of given sediment size class (m³); $MassBed_{Sed}$ mass of the given sediment size class (kg); and density of given sediment size class (kg/m³). Rho_{Sed}

The concentrations of silt and clay in the water column are computed similarly to the mass of those sediments in the bed, with the addition of loadings from upstream and losses downstream:

$$Conc_{Sed} = \frac{KgLoad_{Sed}}{Q \cdot 86400} + Conc_{Sed, t=-1} + Scour_{Sed} - Deposit_{Sed} - Wash_{Sed}$$
 (159)

where:

concentration of silt or clay in water column (kg/m³); $Conc_{Sed}$ $Conc_{Sed, t = -1}$ concentration of silt or clay on previous day (kg/m³);

 $KgLoad_{Sed}$ loading of clay or silt (kg/d);

flow rate (m^3/s converted to m^3/d);

 $Scour_{Sed}$ $Deposit_{Sed}$ amount of silt or clay resuspended (kg/m³);

amount of suspended sediment deposited (kg/m³); and

 $Wash_{Sed}$ amount of sediment lost through downstream transport (kg/m³).

The concentration of sand is computed using a totally different approach, which is described in Section 6.2.

6.1 Deposition and Scour of Silt and Clay

Relationships for scour and deposition of cohesive sediments (silts and clays) used in AQUATOX are the same as the ones used by the Hydrologic Simulation Program in Fortran (HSPF, US EPA 1991). Deposition and scour of silts and clay are modeled using the relationships for deposition (Krone, 1962) and scour (Partheniades, 1965) as summarized by Partheniades (1971).

Shear stress is computed as (Bicknell et al., 1992):

$$Tau = H2ODensity \cdot Slope \cdot HRadius$$
 (160)

where:

Tau = shear stress (kg/m^2) ;

density of water (1000 kg/m³); H2ODensity =slope of channel (m/m); Slope

and hydraulic radius (*HRadius*) is (Colby and McIntire, 1978):

$$HRadius = \frac{Y \cdot Width}{2 \cdot Y + Width}$$
 (161)

where:

hydraulic radius (m);

HRadius = Y = dynamic mean depth (m); and

Width channel width (m).

Resuspension or scour of bed sediments is predicted to occur when the computed shear stress is greater than the critical shear stress for scour:

$$Scour_{Sed} = \frac{Erodibility_{Sed}}{Y} \cdot \left(\frac{Tau}{TauScour_{Sed}} - 1\right)$$
(162)

where:

 $Scour_{Sed} = Erodibility_{Sed} = TauScour_{Sed} =$ resuspension of silt or clay (kg/m³); erodibility coefficient (kg/m²); and

 $TauScour_{Sed} =$ critical shear stress for scour of silt or clay (kg/m²).

The amount of sediment that is resuspended is constrained by the mass of sediments stored in the bed. An intermediate variable representing the maximum potential mass that can be scoured is calculated; if the mass available is less than the potential, then scour is set to the lower amount:

$$Check_{Sed} = Scour_{Sed} \cdot Volume_{Water}$$
 (163)

if
$$Mass_{Sed} \leq Check_{Sed}$$
 then
$$Scour_{Sed} = \frac{Mass_{Sed}}{Volume_{Water}}$$
(164)

where:

 $Check_{Sed} = Mass_{Sed} =$ maximum potential mass (kg); and mass of silt or clay in bed (kg).

Deposition occurs when the computed shear stress is less than the critical depositional shear stress:

if
$$Tau < TauDep_{Sed}$$
 then
$$Deposit_{Sed} = Conc_{Sed} \cdot \left(1 - \frac{VT_{Sed}}{Y} \cdot \left(1 - \frac{Tau}{TauDep_{Sed}}\right)\right)$$
(165)

where:

 $Deposit_{Sed}$ = amount of sediment deposited (kg/m³); $TauDep_{Sed}$ = critical depositional shear stress (kg/m²); $Conc_{Sed}$ = concentration of silt, clay, or sand (kg/m³); and VT_{Sed} = terminal fall velocity of given sediment type (m/s).

The settling velocity is computed from Stoke's law (Schnoor, 1987):

$$VT_{Sed} = \frac{g}{18 \cdot Visc} \cdot \frac{Rho_{Sed} - Rho}{Rho} \cdot \left(\frac{D_{Sed}}{1000}\right)^2$$
 (166)

where:

g = gravitational acceleration constant (9.807 m/s²);

Visc = kinematic viscosity of water (m²/s);

Rho = density of water (kg/m^3) :

 $Rho_{S_{ed}}$ = density of given sediment (kg/m³); and

 D_{Sed} = particle diameter for given sediment (mm, converted to m).

Downstream transport is an important mechanism for loss of suspended sediment from a given stream reach. In a steady-state simulation with constant flow and volume and with a one-day time step, the downstream transport of sediments is simply the amount of sediments in suspension in the previous time step:

$$Wash_{Sed} = Conc_{Sed, t=-1}$$

where:

 $Wash_{Sed}$ = amount of given sediment lost to downstream transport (kg/m³).

6.2 Scour, Deposition and Transport of Sand

Scour, deposition and transport of sand are simulated using the Engelund and Hansen (1967) sediment transport relationships as presented by Brownlie (1981). This relationship was selected

because of its simplicity and accuracy. Brownlie (1981) shows that this relationship gives good results when compared to 13 others using a field and laboratory data set of about 7,000 records.

$$ntConc_{Sand} = 0.05 \cdot \frac{Rho}{Rho_{Sand} - Rho} \cdot \frac{Velocity \cdot Slope}{\sqrt{\frac{Rho_{Sand} - Rho}{Rho} \cdot g \cdot D_{Sand}/1000}} \cdot \sqrt{TauSt}$$
(168)

where:

 $PotConc_{Sand} =$ potential concentration of suspended sand (kg/m³);

 $PotConc_{Sand}$ Rho_{Sand} = Velocity = Slope = =density of sand (kg/m³); flow velocity (m/s); slope of stream (m/m);

mean diameter of sand particle (mm converted to m); and

TauStar dimensionless shear stress.

The flow velocity is calculated by:

$$Velocity = \frac{Q}{Y \cdot Width}$$
 (169)

where:

= flow rate (m³/s); = dynamic mean do

dynamic mean depth of water (m); and

Width channel width (m).

The dimensionless shear stress is calculated by:

$$TauStar = \frac{Rho}{Rho_{Sand} - Rho} \cdot HRadius \cdot \frac{Slope}{D_{Sand}/1000}$$
 (170)

where:

hydraulic radius (m). **HRadius**

Once the potential concentration has been determined for the given flow rate and channel characteristics, it is compared with the present concentration. If the potential concentration is greater, the difference is considered to be made available through scour, up to the limit of the bed. If the potential concentration is less than what is in suspension, the difference is considered to be deposited:

$$Check_{Sand} = PotConc_{Sand} \cdot Volume_{Water}$$
 (171)

$$MassSusp_{Sand} = Conc_{Sand} \cdot Volume_{Water}$$
 (172)

$$TotalMass_{Sand} = MassSusp_{Sand} + MassBed_{Sand}$$
 (173)

if
$$Check_{Sand} \leq MassSusp_{Sand}$$
 then

$$Deposit_{Sand} = MassSusp_{Sand} - Check_{Sand}$$

$$Conc_{Sand} = PotConc_{Sand}$$
(174)

if
$$Check_{Sand} \geq TotalMass_{Sand}$$
 then
$$MassBed_{Sand} = 0$$

$$Conc_{Sand} = \frac{TotalMass_{Sand}}{Volume_{Water}}$$
(175)

if
$$Check_{Sand} > MassSusp_{Sand}$$
 and $< TotalMass_{Sand}$ then
$$Scour_{Sand} = Check_{Sand} - MassSusp_{Sand}$$

$$Conc_{Sand} = \frac{MassSusp_{Sand} + Scour_{Sand}}{Volume_{Weatorn}}$$
(176)

6.3 Suspended Inorganic Sediments in Standing Water

At present, AQUATOX does not compute settling of inorganic sediments in standing water or scour as a function of wave action. However, suspended sediments are important in creating turbidity and limiting light, especially in reservoirs and shallow lakes. Therefore, the user can provide loadings of total suspended solids (TSS), and the model will back-calculate suspended inorganic sediment concentrations by subtracting predicted phytoplankton and suspended detritus concentrations:

$$InorgSed = TSS - \sum Phyto - \sum PartDetr$$
 (177)

InorgSed=concentration of suspended inorganic sediments (g/m³);TSS=observed concentration of total suspended solids (g/m³);Phyto=predicted phytoplankton concentrations (g/m³); see (25) and

PartDetr = predicted suspended detritus concentrations (g/m³); see (94) and (95).

The concentration of suspended inorganic sediments is used solely to calculate their contribution to the extinction coefficient, which affects the depth of the euphotic zone and the Secchi depth (see (30)).

7. TOXIC ORGANIC CHEMICALS

The chemical fate module of AQUATOX predicts the partitioning of a compound between water, sediment, and biota (**Figure 65**), and estimates the rate of degradation of the compound (**Figure 66**). Microbial degradation, photolysis, hydrolysis, and volatilization are modeled in AQUATOX. Each of these processes is described generally, and again in more detail below.

Nonequilibrium concentrations, as represented by kinetic equations, depend on sorption, desorption, and elimination as functions of the chemical and exposure through water and food as a function of bioenergetics of the organism. Equilibrium partitioning is no longer represented in AQUATOX.

Microbial degradation is modeled by entering a maximum biodegradation rate for a particular organic toxicant, which is subsequently reduced to account for suboptimal temperature, pH, and dissolved oxygen. Photolysis is modeled by using a light screening factor (Schwarzenbach et al., 1993) and the near-surface, direct photolysis first-order rate constant for each pollutant. The light screening factor is a function of both the diffuse attenuation coefficient near the surface and the average diffuse attenuation coefficient for the whole water column. For those organic chemicals that undergo hydrolysis, neutral, acid-, and base-catalyzed reaction rates are entered into AQUATOX as applicable. Volatilization is modeled using a stagnant two-film model, with the air and water transfer velocities approximated by empirical equations based on reaeration of oxygen (Schwarzenbach et al., 1993).

Figure 65In-situ Uptake and Release of Insecticide

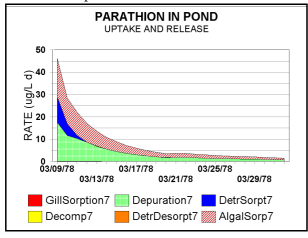
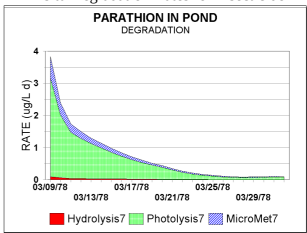


Figure 66In-situ Degradation Rates for Insecticide



The mass balance equations follow. The change in mass of toxicant in the water includes explicit representations of mobilization of the toxicant from sediment to water as a result of decomposition of the labile sediment detritus compartment, sorption to and desorption from the detrital sediment compartments, uptake by algae and macrophytes, uptake across the gills of animals, depuration by organisms, and turbulent diffusion between epilimnion and hypolimnion:

$$\frac{\text{d}Toxicant_{Water}}{\text{d}t} = Loading + \sum_{LabileDetr} (Decomposition_{LabileDetr} \cdot PPB_{LabileDetr} \cdot 1e-6) \\ + \sum_{Desorption_{DetrTox}} + \sum_{Org} (K2 \cdot PPB_{Org} \cdot 1e-6) \\ - \sum_{Sorption_{DetrTox}} - \sum_{GillUptake_{Pred}} - MacroUptake \\ - \sum_{AlgalUptake_{Alga}} - Hydrolysis - Photolysis - MicrobialDegrdn \\ - Volatilization - Discharge + TurbDiff$$

$$(178)$$

The equations for the toxicant associated with the two sediment detritus compartments are rather involved:

$$\frac{\text{d} Toxicant_{SedLabileDetr}}{\text{d} t} = Sorption - Desorption \\ + Colonization \cdot PPB_{SedRefrDetr} \cdot 1 e - 6 \\ + \sum_{Pred} \sum_{Prey} \left(Def2Detr \cdot DefecationTox_{Pred, Prey} \right) \\ - (Resuspension + Decomposition) \cdot PPB_{SedLabileDetr} \cdot 1 e - 6 \\ - \sum_{Pred} Ingestion_{Pred, SedLabileDetr}) \cdot PPB_{SedLabileDetr} \cdot 1 e - 6 \\ + Sedimentation \cdot PPB_{SuspLabileDetr} \cdot 1 e - 6 \\ + \sum_{Pred} (Sed2Detr \cdot Sink_{Phyto} \cdot PPB_{Phyto} \cdot 1 e - 6) \\ - Hydrolysis - MicrobialDegran - Burial + Expose$$
 (179)

$$\frac{\text{d}Toxicant_{SedRefrDetr}}{\text{d}t} = Sorption - Desorption \\ + \sum_{Pred} \sum_{Prey} \left((1 - Def2Detr) \cdot DefecationTox_{Pred, Prey} \right) \\ - (Resuspension + Colonization) \cdot PPB_{SedRefrDetr} \cdot 1e - 6 \\ - \sum_{Pred} Ingestion_{Pred, SedRefrDetr}) \cdot PPB_{SedRefrDetr} \cdot 1e - 6 \\ + Sedimentation \cdot PPB_{SuspRefrDetr} \cdot 1e - 6 \\ + \sum (Sed2Detr \cdot Sink_{Phyto} \cdot PPB_{Phyto} \cdot 1e - 6) \\ - Hydrolysis - MicrobialDegran - Burial + Expose$$
 (180)

Similarly for the toxicant associated with suspended and dissolved detritus, the equations are:

$$\frac{Toxicant_{SuspLabileDetr}}{\mathrm{d}t} = Loading + Sorption - Desorption + \sum_{Pred} (Def2Sed \cdot Def_{Pred})$$

$$+ \sum_{Org} ((Mort2Detr \cdot Mortality_{Org} + GameteLoss_{Org})$$

$$\cdot PPB_{Org} \cdot 1e - 6) - (Sedimentation + Washout + Decomp$$

$$+ \sum_{Pred} Ingestion_{Pred, SuspLabileDetr}) \cdot PPB_{SuspLabileDetr} \cdot 1e - 6$$

$$+ Colonization \cdot PPB_{SuspRefrDetr} \cdot 1e - 6$$

$$+ Resuspension \cdot PPB_{SedLabileDetr} \cdot 1e - 6 - SedToHyp + SedFrE_{I}$$

$$- Hydrolysis - Photolysis - MicrobialDegran + TurbDiff$$

$$\frac{\text{d} Toxicant_{SuspRefrDetr}}{\text{d} t} = Loading + Sorption - Desorption \\ + \sum_{Org} (Mort2Ref \cdot Mortality_{Org} \cdot PPB_{Org} \cdot 1e - 6) \\ - (Sedimentation + Washout + Colonization \\ + \sum_{Pred} Ingestion_{SuspRefrDetr}) \cdot PPB_{SuspRefrDetr} \cdot 1e - 6 \\ + Resuspension \cdot PPB_{SedRefrDetr} \cdot 1e - 6 - SedToHyp + SedFrEpi \\ - Hydrolysis - Photolysis - MicrobialDegran + TurbDiff$$

$$\frac{\text{d}Toxicant_{DissLabileDetr}}{\text{d}t} = Loading + Sorption - Desorption + \sum ExcrToxToDiss_{Org} \\ + \sum_{Org} (Mort2Detr \cdot Mortality_{Org} \cdot PPB_{Org} \cdot 1e - 6) \\ - (Washout + Decomposition) \cdot PPB_{DissLabileDetr} \cdot 1e - 6 \\ + Colonization \cdot PPB_{DissRefrDetr} \cdot 1e - 6 \\ - Hydrolysis - Photolysis - MicrobialDegran + TurbDiff$$
 (183)

$$\frac{\text{d}Toxicant_{DissRefrDetr}}{\text{d}t} = Loading + Sorption - Desorption + \sum ExcToxToDiss_{Org} \\ + \sum_{Org} (Mort2Ref \cdot Mortality_{Org} \cdot PPB_{Org} \cdot 1 \text{ e} - 6) \\ - (Washout + Colonization) \cdot PPB_{DissRefrDetr} \cdot 1 \text{ e} - 6 \\ - Hydrolysis - Photolysis - MicrobialDegran + TurbDiff}$$

$$(184)$$

Note that there are no equations for buried detritus, as they are considered to be sequestered and outside of the influence of any processes which would change the concentrations of their associated toxicants.

Algae are represented as:

$$\frac{dToxicant_{Alga}}{dt} = Loading + AlgalUptake - Depuration + TurbDiff + (- Excretion - Washout - $\sum_{Pred} Predation_{Pred, Alga} - Mortality - Sink + SinkToHypo - SinkFrEpi) \cdot PPB_{Alga} \cdot 1e-6$ (185)$$

Macrophytes are represented similarly, but reflecting the fact that they are stationary:

$$\frac{\mathrm{d}Toxicant_{Macrophyte}}{\mathrm{d}t} = Loading + MacroUptake - Depuration - (Excretion + \sum_{Pred} Predation_{Pred, Macro} + Mortality) \cdot PPB_{Macro} \cdot 1e - 6$$
(186)

The toxicant associated with animals is represented by an involved kinetic equation because of the various routes of exposure and transfer:

$$\frac{\text{d}Toxicant_{Animal}}{\text{d}t} = Loading + GillUptake + \sum_{Prey} DietUptake + TurbDiff$$

$$- (Depuration + \sum_{Pred} Predation_{Pred, Animal} + Mortality + Spawn$$

$$\pm Promotion + Drift + Migration + EmergeInsect) \cdot PPB_{Animal} \cdot 1e-6$$
(187)

where:

 $Toxicant_{Water}$ toxicant in dissolved phase in unit volume of water ($\mu g/L$); = $Toxicant_{SedDetr}$ mass of toxicant associated with each of the two sediment detritus compartments in unit volume of water (µg/L); mass of toxicant associated with each of the two suspended $Toxicant_{SuspDetr}$ = detritus compartments in unit volume of water (µg/L); $Toxicant_{DissDetr}$ mass of toxicant associated with each of the two dissolved = organic compartments in unit volume of water ($\mu g/L$); mass of toxicant associated with given alga in unit volume of $Toxicant_{Alog}$ =water (μ g/L); $Toxicant_{Macrophyte}$ mass of toxicant associated with macrophyte in unit volume =of water(μ g/L); $Toxicant_{Animal}$ mass of toxicant associated with given animal in unit volume = of water ($\mu g/L$); $PPB_{SedDetr}$ concentration of toxicant in sediment detritus (µg/kg); = $PPB_{SuspDetr}$ = concentration of toxicant in suspended detritus (µg/kg);

 $PPB_{DissDetr}$ = concentration of toxicant in dissolved organics ($\mu g/kg$);

 PPB_{Alga} = concentration of toxicant in given alga (μ g/kg); $PPB_{Macrophyte}$ = concentration of toxicant in macrophyte (μ g/kg); PPB_{Animal} = concentration of toxicant in given animal (μ g/kg);

1 e -6 = units conversion (kg/mg);

Loading = loading of toxicant from external sources (μ g/L·d);

TurbDiff = depth-averaged turbulent diffusion between epilimnion and

hypolimnion (µg/L·d), see 11;

Hydrolysis = rate of loss due to hydrolysis (μ g/L·d), see (190);

Photolysis = rate of loss due to direct photolysis (μ g/L·d), see (197);

MicrobialDegrdn = rate of loss due to microbial degradation ($\mu g/L \cdot d$), see (204);

Volatilization = rate of loss due to volatilization ($\mu g/L \cdot d$), see (209);

Discharge = rate of loss of toxicant due to discharge downstream (μ g/L·d),

see Table 1;

Burial = rate of loss due to deep burial (μ g/L·d) see (165);

Expose = rate of exposure due to resuspension of overlying sediments

 $(\mu g/L \cdot d)$, see (162);

Decomposition = rate of decomposition of given detritus (mg/L·d), see (112);

Depuration = elimination rate for toxicant due to clearance (μ g/L·d), see

(258);

Sorption = rate of sorption to given compartment (μ g/L·d), see (230);

Desorption = rate of desorption from given compartment ($\mu g/L \cdot d$), see

(231);

Colonization = rate of conversion of refractory to labile detritus (g/m³·d), see

(108);

 $DefecationTox_{Pred, Prev}$ = rate of transfer of toxicant due to defecation of given prey by

given predator (μ g/L·d), see (259);

Def2Detr=fraction of defecation that goes to given compartment;Resuspension=rate of resuspension of given sediment detritus (mg/L·d);Sedimentation=rate of sedimentation of given suspended detritus (mg/L·d);Sed2Detr=fraction of sinking phytoplankton that goes to given detrital

compartment;

Sink = loss rate of phytoplankton to bottom sediments (mg/L·d), see

(57);

Death = nonpredatory mortality of given organism (mg/L·d), see (80);

Mort2Detr = fraction of dead organism that is labile (unitless);

GameteLoss = loss rate for gametes $(g/m^3 \cdot d)$, see (84);

Mort2Ref = fraction of dead organism that is refractory (unitless);

Washout or Drift = rate of loss of given suspended detritus or organism due to

being carried downstream (mg/L·d), see (13);

SedToHyp = rate of settling loss to hypolimnion from epilimnion (mg/L·d);

SedFrEpi = rate of gain to hypolimnion from settling out of epilimnion

 $(mg/L\cdot d);$

 $Ingestion_{Pred, Prey}$ = rate of ingestion of given food or prey by given predator

 $(mg/L\cdot d)$, see (68);

 $Predation_{Pred, Prev}$ = predatory mortality by given predator on given prey (mg/L·d),

see (75);

 $ExcToxToDiss_{Org}$ = toxicant excretion from plants to dissolved organics (mg/L·d);

Excretion = excretion rate for given organism $(g/m^3 \cdot d)$, see (79);

SinkToHypo = rate of transfer of phytoplankton to hypolimnion (mg/L·d);

SinkFrEpi = loss rate of phytoplankton to hypolimnion (mg/L·d);

AlgalUptake = rate of sorption by algae (μ g/L - d), see (244);

MacroUptake = rate of sorption by macrophytes (µg/L - d), see (240);

GillUptake = rate of absorption of toxicant by the gills (μ g/L - d), see (249); DietUptake_{Prev} = rate of dietary absorption of toxicant associated with given

prey (µg/L·d), see (252);

Promotion = promotion from one age class to the next (mg/L·d), see (92);

Migration = rate of migration (g/m 3 ·d), see (91); and EmergeInsect = insect emergence (mg/L·d), see (93).

7.1 Ionization

Dissociation of an organic acid or base in water can have a significant effect on its environmental properties. In particular, solubility, volatilization, photolysis, sorption, and bioconcentration of an ionized compound can be affected. Rather than modeling ionization products, the approach taken in AQUATOX is to represent the modifications to the fate and transport of the neutral species, based on the fraction that is not dissociated. The acid dissociation constant is expressed as the negative \log , pKa, and the fraction that is not ionized is:

$$Nondissoc = \frac{1}{1 + 10^{(pH - pKa)}}$$
 (188)

where:

Nondissoc = nondissociated fraction (unitless).

If the compound is a base then the fraction not ionized is:

$$Nondissoc = \frac{1}{1 + 10^{(pKa - pH)}}$$
 (189)

When pKa = pH half the compound is ionized and half is not (**Figure 67**). At ambient environmental pH values, compounds with a pKa in the range of 4 to 9 will exhibit significant dissociation (**Figure 68**).

Figure 67
Dissociation of Pentachlorophenol (pKa = 4.75) at Higher pH Values

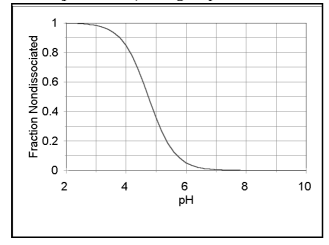
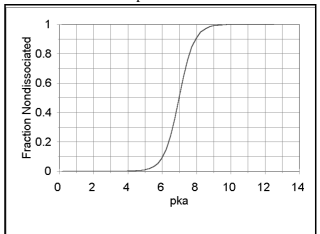


Figure 68Dissociation as a Function of *pKa* at an Ambient pH of 7



7.2 Hydrolysis

Hydrolysis is the degradation of a compound through reaction with water. During hydrolysis, both a pollutant molecule and a water molecule are split, and the two water molecule fragments (H⁺ and OH⁻) join to the two pollutant fragments to form new chemicals. Neutral and acid- and base-catalyzed hydrolysis are modelled using the approach of Mabey and Mill (1978) in which an overall pseudo-first-order rate constant is computed for a given pH, adjusted for the ambient temperature of the water:

$$Hydrolysis = KHyd \cdot Toxicant_{Phase}$$
 (190)

where:

$$KHyd = (KAcidExp + KBaseExp + KUncat) \cdot Arrhen$$
 (191)

and where:

KHyd = overall pseudo-first-order rate constant for a given pH and temperature

(1/d);

KAcidExp = pseudo-first-order acid-catalyzed rate constant for a given pH (1/d); *KBaseExp* = pseudo-first-order base-catalyzed rate constant for a given pH (1/d);

KUncat = the measured first-order reaction rate at pH 7 (1/d); and

Arrhen = temperature adjustment (unitless).

There are three types of hydrolysis: acid-catalyzed, base-catalyzed, and neutral. In neutral hydrolysis reactions, the pollutant reacts with a water molecule (H_2O) and the concentration of water is usually included in KUncat. In acid-catalyzed hydrolysis, the hydrogen ion reacts with the pollutant, and a first-order decay rate for a given pH can be estimated as follows:

$$KAcidExp = KAcid \cdot HIon$$
 (192)

where:

$$HIon = 10^{-pH} \tag{193}$$

and where:

KAcid = acid-catalyzed rate constant (L/mol·d);
HIon = concentration of hydrogen ions (mol/L); and

pH = pH of water column.

Likewise for base-catalyzed hydrolysis, the first-order rate constant for a reaction between the hydroxide ion and the pollutant at a given pH (**Figure 69**) can be described as:

$$KBaseExp = KBase \cdot OHIon$$
 (194)

where:

$$OHIon = 10^{pH - 14}$$
 (195)

and where:

KBase = base-catalyzed rate constant (L/mol·d); and
 OHIon = concentration of hydroxide ions (mol/L).

6.05E-03 6.00E-03 EDITION OF THE INTERPOLATION OF

Figure 69Base-catalyzed Hydrolysis of Pentachlorophenol

Hydrolysis reaction rates were adjusted for the temperature of the waterbody being modeled by using the Arrhenius rate law (Hemond and Fechner, 1994). An activation energy value of 18,000 cal/mol (a mid-range value for organic chemicals) was used as a default:

$$Arrhen = e^{-\left(\frac{En}{R \cdot KelvinT} - \frac{En}{R \cdot TObs}\right)}$$
 (196)

where:

En = Arrhenius activation energy (cal/mol); R = universal gas constant (cal/mol · Kelvin);

KelvinT = temperature for which rate constant is to be predicted (Kelvin); and *TObs* = temperature at which known rate constant was measured (Kelvin).

7.3 Photolysis

Direct photolysis is the process by which a compound absorbs light and undergoes transformation:

$$Photolysis = KPhot \cdot Toxicant_{Phase}$$
 (197)

where:

Photolysis = rate of loss due to photodegradation $(g/m^3 \cdot d)$; and KPhot = direct photolysis first-order rate constant (1/day).

For consistency, photolysis is computed for both the epilimnion and hypolimnion in stratified systems. However, it is not a significant factor at hypolimnetic depths.

Ionization may result in a significant shift in the absorption of light (Lyman et al., 1982; Schwarzenbach et al., 1993). However, there is a general absence of information on the effects of light on ionized species. The user provides an observed half-life for photolysis, and this is usually determined either with distilled water or with water from a representative site, so that ionization may be included in the calculated lumped parameter *KPhot*.

Based on the approach of Thomann and Mueller (1987; see also Schwarzenbach et al. 1993), the observed first-order rate constant for the compound is modified by a light attenuation factor for ultraviolet light so that the process as represented is depth-sensitive (**Figure 70**); it also is adjusted by a factor for time-varying light:

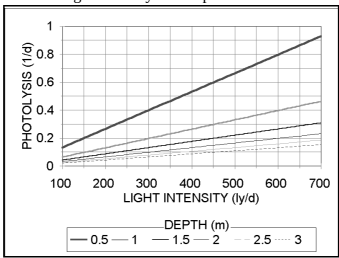
$$KPhot = PhotRate \cdot ScreeningFactor \cdot LightFactor$$
 (198)

where:

PhotRate = direct, observed photolysis first-order rate constant (1/day);

ScreeningFactor = a light screening factor (unitless), see (199); and LightFactor = a time-varying light factor (unitless), see (201).

Figure 70
Photolysis of Pentachlorophenol as a Function of
Light Intensity and Depth of Water



A light screening factor adjusts the observed laboratory photolytic transformation rate of a given pollutant for field conditions with variable light attenuation and depth (Thomann and Mueller, 1987):

$$ScreeningFactor = \frac{RadDistr}{RadDistr0} \cdot \frac{1 - \exp^{(-Alpha \cdot Thick)}}{Alpha \cdot Thick}$$
(199)

RadDistr = radiance distribution function, which is the ratio of the average

pathlength to the depth (see Schwarzenbach et al., 1993) (taken to be

1.6, unitless);

RadDistr0 = radiance distribution function for the top of the segment (taken to be 1.2)

for the top of the epilimnion and 1.6 for the top of the hypolimnion,

unitless);

Alpha = light extinction coefficient at wavelength 312.5 nm (1/m), see (200);

and

Thick = thickness of the water body segment if stratified or maximum depth if

unstratified (m).

The extinction of light of the reference wavelength of 312.5 nm is a function of several components, based on parameter values in Burns and Cline (1985), as given in Ambrose et al. (1991):

$$Alpha = ExtinctH2O + AttenChl \cdot Chlorophyll + AttenDOC \cdot DOC + AttenSolids \cdot SuspSed$$
 (200)

where:

Extinct H2O = light extinction of wavelength 312.5 nm in pure water (1/m);

AttenChl = attenuation coefficient for chlorophyll a (L/mg - m);

AttenDOC = attenuation coefficient for dissolved organic carbon (L/mg - m);
AttenSolids = attenuation coefficient for suspended sediment (L/mg - m);
Chlorophyll = concentration of chlorophyll a in water column (mg/L);

DOC = concentration of dissolved organic carbon in water (mg/L); and

SuspSed = concentration of suspended sediments (mg/L).

The equations presented above implicitly make the following assumptions:

the compound of interest absorbs light only over a relatively narrow wavelength range, and the screening factor for wavelength 312.5 nm can be considered representative;

· quantum yield is independent of wavelength; and,

• the value used for *PhotRate* is a representative near-surface, first-order rate constant for direct photolysis.

The rate is modified further to represent seasonally varying light conditions and the effect of ice cover:

$$LightFactor = \frac{Solar0}{AveSolar}$$
 (201)

where:

Solar0 = time-varying average light intensity at the top of the segment (ly/day);

and

AveSolar = average light intensity for late spring or early summer, corresponding

to time when photolytic half-life is often measured (default = 500

Ly/day).

If the system is unstratified or if the epilimnion is being modeled, the light intensity is the light loading:

$$Solar0 = Solar$$
 (202)

otherwise we are interested in the intensity at the top of the hypolimnion and the attenuation of light is given as a logarithmic decrease over the thickness of the epilimnion:

$$Solar0 = Solar \cdot \exp^{(-Alpha \cdot MaxZMix)}$$
 (203)

where:

Solar = incident solar radiation loading (ly/d), see (22); and

MaxZMix = depth of the mixing zone (m), see (14).

Because the ultraviolet light intensity exhibits greater seasonal variation than the visible spectrum (Lyman et al., 1982), decreasing markedly when the angle of the sun is low, this construct could predict higher rates of photolysis in the winter than might actually occur. However, the model also accounts for significant attenuation of light due to ice cover so that photolysis, as modeled, is not an important process in northern waters in the winter.

7.4 Microbial Degradation

Not only can microorganisms decompose the detrital organic material in ecosystems, they also can degrade xenobiotic organic compounds such as fuels, solvents, and pesticides to obtain energy. In AQUATOX this process of biodegradation of pollutants, whether they are dissolved in the water column or adsorbed to organic detritus in the water column or sediments, is modeled using the same equations as for decomposition of detritus, substituting the pollutant and its degradation parameters for detritus in Equation (112) and supporting equations:

$$\label{eq:microbialDegrdn} \begin{aligned} \textit{MicrobialDegrdn} &= \textit{KMDegrdn}_{\textit{Phase}} \cdot \textit{DOCorrection} \cdot \textit{TCorr} \cdot \textit{pHCorr} \\ &\cdot \textit{Toxicant}_{\textit{Phase}} \end{aligned} \tag{204}$$

where:

MicrobialDegrdn = loss due to microbial degradation (g/m 3 ·d);

KMDegrdn = maximum degradation rate, either in water column or sediments

(1/d);

DOCorrection=effect of anaerobic conditions (unitless), see (113);TCorr=effect of suboptimal temperature (unitless), see (21);pHCorr=effect of suboptimal pH (unitless), see (115); and

Toxicant = concentration of organic toxicant (g/m^3) .

Microbial degradation proceeds more quickly if the material is associated with surficial sediments rather than suspended in the water column (Godshalk and Barko, 1985); thus, in calculating the loss due to microbial degradation in the dissolved phase, the maximum degradation rate is set arbitrarily to 25 percent of the maximum degradation rate in the sediments. The model assumes that reported maximum microbial degradation rates are for suspended slurry or wet soil samples; if the reported degradation value is from a flask study without additional organic matter, then the parameter value that is entered should be four times that reported.

7.5 Volatilization

Volatilization is modeled using the "stagnant boundary theory", or two-film model, in which a pollutant molecule must diffuse across both a stagnant water layer and a stagnant air layer to volatilize out of a waterbody (Whitman, 1923; Liss and Slater, 1974). Diffusion rates of pollutants in these stagnant boundary layers can be related to the known diffusion rates of chemicals such as oxygen and water vapor. The thickness of the stagnant boundary layers must also be taken into account to estimate the volatile flux of a chemical out of (or into) the waterbody.

The time required for a pollutant to diffuse through the stagnant water layer in a waterbody is based on the well-established equations for the reaeration of oxygen, corrected for the difference in diffusivity as indicated by the respective molecular weights (Thomann and Mueller, 1987, p. 533). The diffusivity through the water film is greatly enhanced by the degree of ionization (Schwarzenbach et al., 1993, p. 243), and the depth-averaged reaeration coefficient is multiplied by the thickness of the well-mixed zone:

$$KLiq = KReaer \cdot Thick \cdot \left(\frac{MolWtO2}{MolWt}\right)^{0.25} \cdot \frac{1}{Nondissoc}$$
 (205)

where:

KLiq = water-side transfer velocity (m/d);

KReaer = depth-averaged reaeration coefficient for oxygen (1/d), see (139)-(146); Thick = thickness of the water body segment if stratified or maximum depth if

unstratified (m);

MolWtO2=molecular weight of oxygen (g/mol, =32);MolWt=molecular weight of pollutant (g/mol); andNondissoc=nondissociated fraction (unitless), see (188).

Likewise, the thickness of the air-side stagnant boundary layer is also affected by wind. To estimate the air-side transfer velocity of a pollutant, we used the following empirical equation based on the evaporation of water, corrected for the difference in diffusivity of water vapor compared to the toxicant (Thomann and Mueller, 1987, p. 534):

$$KGas = 168 \cdot \left(\frac{MolWtH2O}{MolWt}\right)^{0.25} \cdot Wind$$
 (206)

where:

KGas = air-side transfer velocity (m/d);

Wind = wind speed ten meters above the water surface (m/s); and

MolWtH2O = molecular weight of water (g/mol, =18).

The total resistance to the mass transfer of the pollutant through both the stagnant boundary layers can be expressed as the sum of the resistances—the reciprocals of the air- and water-phase mass transfer coefficients (Schwarzenbach et al., 1993), modified for the effects of ionization:

$$\frac{1}{KOVol} = \frac{1}{KLiq} + \frac{1}{KGas \cdot HenryLaw \cdot Nondissoc}$$
 (207)

where:

KOVol = total mass transfer coefficient through both stagnant boundary layers (m/d);

$$HenryLaw = \frac{Henry}{R \cdot TKelvin}$$
 (208)

and where:

HenryLaw = Henry's law constant (unitless); Henry = Henry's law constant (atm m³ mol⁻¹);

R = gas constant (=8.206E-5 atm m³ (mol K)⁻¹); and

TKelvin temperature in °K.

The Henry's law constant is applicable only to the fraction that is nondissociated because the ionized species will not be present in the gas phase (Schwarzenbach et al., 1993, p. 179).

The atmospheric exchange of the pollutant can be expressed as the depth-averaged total mass transfer coefficient times the difference between the concentration of the chemical and the saturation concentration:

$$Volatilization = \frac{KOVol}{Thick} \cdot \left(ToxSat - Toxicant_{water} \right)$$
 (209)

where:

Volatilization = interchange with atmosphere ($\mu g/L \cdot d$);

Thick depth of water or thickness of surface layer (m); saturation concentration of pollutant (µg/L); and **ToxSat** Toxicant_{water} concentration of pollutant in water (µg/L).

The saturation concentration depends on the concentration of the pollutant in the air, ignoring temperature effects (Thomann and Mueller, 1987, p. 532), but adjusting for ionization and units:

$$ToxSat = \frac{Toxicant_{air}}{HenryLaw \cdot Nondissoc} \cdot 1000$$
 (210)

where:

 $Toxicant_{air} = Nondissoc =$ gas-phase concentration of the pollutant (g/m³); and

Nondissoc nondissociated fraction (unitless).

Often the pollutant can be assumed to have a negligible concentration in the air and *ToxSat* is zero. However, this general construct can represent the transferral of volatile pollutants into water bodies. Because ionized species do not volatilize, the saturation level increases if ionization is occurring.

The nondimensional Henry' saw constant, which relates the concentration of a compound in the air phase to its concentration in the water phase, strongly affects the air-phase resistance. Depending on the value of the Henry' kaw constant, the water phase, the air phase or both may control volatilization. For example, with a depth of 1 m and a wind of 1 m/s, the gas phase is 100,000 times as important as the water phase for atrazine (Henry' kaw constant = 3.0E-9), but the water phase is 50 times as important as the air phase for benzene (Henry's law constant = 5.5E-3). Volatilization of atrazine exhibits a linear relationship with wind (Figure 71) in contrast to the exponential relationship exhibited by benzene (Figure 72).

Figure 71 Atrazine *KOVol* as a Function of Wind

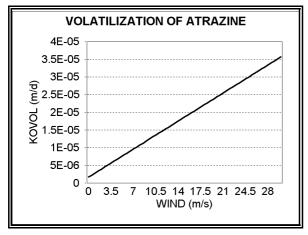
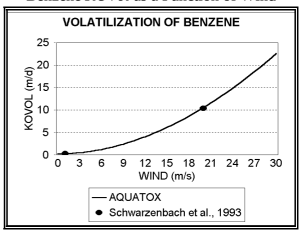


Figure 72 Benzene *KOVol* as a Function of Wind



7.6 Partition Coefficients

Although AQUATOX is a kinetic model, steady-state partition coefficients for organic pollutants are computed in order to place constraints on competitive uptake and loss processes, speeding up computations. They are estimated from empirical regression equations and the pollutant's octanol-water partition coefficient.

Natural organic matter is the primary sorbent for neutral organic pollutants. Hydrophobic chemicals partition primarily in nonpolar organic matter (Abbott et al. 1995). Refractory detritus is relatively nonpolar; its partition coefficient is a function of the octanol-water partition coefficient (N = 34, $r^2 = 0.93$; Schwarzenbach et al. 1993):

$$KOM_{RefrDetr} = 1.38 \cdot KOW^{0.82} \tag{211}$$

where:

 $KOM_{RefrDetr}$ = detritus-water partition coefficient (L/kg); and KOW = octanol-water partition coefficient (unitless).

This and the following equations are extended to polar compounds, following the work of Smejtek and Wang (1993):

$$KOM_{RefrDetr} = 1.38 \cdot KOW^{0.82} \cdot Nondissoc$$

+ $(1 - Nondissoc) \cdot IonCorr \cdot 1.38 \cdot KOW^{0.82}$ (212)

where:

Nondissoc = un-ionized fraction (unitless); and

IonCorr = correction factor for decreased sorption, generally 0.1 (unitless).

Using pentachlorophenol as a test compound, and comparing it to octanol, the influence of pH-mediated dissociation is seen in **Figure 73**. This relationship is verified by comparison with the results of Smejtek and Wang (1993) using egg membrane.

Partitioning of bioaccumulative chemicals on organic carbon in sediments in Lake Ontario, as represented by the Oliver and Niimi (1988) data, exhibits a weak relationship with KOW (US EPA 1995, Burkhard 1998):

$$KOC = 25 \cdot KOW$$
 (213)

where:

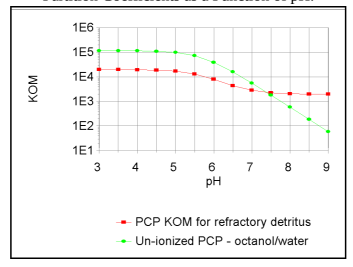
KOC = the partition coefficient for particulate organic carbon.

Converting to organic matter and generalizing to include polar compounds, this relationship was used in AQUATOX to represent the partitioning of chemicals between water and refractory detritus in sediments in a validation for Lake Ontario (Park, 1999c, in Volume 3):

$$KOM = 13 \cdot KOW + (1 - Nondissoc) \cdot IonCorr \cdot 13 \cdot KOW$$
 (214)

However, in the general model Eq. (212) is used for refractory detrital sediments as well.

Figure 73
Refractory Detritus-water and Octanol-water
Partition Coefficients as a Function of pH.



There appears to be a dichotomy in partitioning; data in the literature suggest that labile detritus does not take up hydrophobic compounds as rapidly as refractory detritus. Algal cell membranes contain polar lipids, and it is likely that this polarity is retained in the early stages of decomposition. KOC does not remain the same upon aging, death, and decomposition, probably because of polarity changes. In an experiment using fresh and aged algal detritus, there was a 100% increase in KOC with aging (Koelmans et al., 1995). KOC increased as the C/N ratio increased, indicating that the material was becoming more refractory. In another study, KOC doubled between day 2 and day 34, probably due to deeper penetration into the organic matrix and lower polarity (Cornelissen et al., 1997).

Polar substrates increase the pKa of the compound (Smejtek and Wang, 1993). This is represented in the model by lowering the pH of polar particulate material by one pH unit, which changes the dissociation accordingly.

The partition equation for labile detritus (N = 3, $r^2 = 1.0$;) is based on a study by Koelmans et al. (1995) using fresh algal detritus:

$$KOC_{LahPart} = 23.44 \cdot KOW^{0.61}$$
 (215)

The equation is generalized to polar compounds and transformed to an organic matter partition coefficient:

$$KOM_{LabDetr} = (23.44 \cdot KOW^{0.61} \cdot Nondissoc + (1 - Nondissoc) \cdot IonCorr \cdot 23.44 \cdot KOW^{0.61}) \cdot 0.526$$
(216)

where:

 $KOC_{LabPart}$ = partition coefficient for labile particulate organic carbon (L/kg); and $KOM_{LabDetr}$ = partition coefficient for labile detritus (L/kg).

O'Connor and Connolly (1980; see also Ambrose et al., 1991) found that the sediment partition coefficient is the inverse of the mass of suspended sediment, and Di Toro (1985) developed a construct to represent the relationship. However, AQUATOX models partitioning directly to organic detritus and ignores inorganic sediments, which are seldom involved directly in sorption of neutral organic pollutants. Therefore, the partition coefficient is not corrected for mass of sediment.

Association of hydrophobic compounds with colloidal and dissolved organic matter (DOM) reduces bioavailability; such contaminants are unavailable for uptake by organisms (Stange and Swackhamer 1994, Gilek et al. 1996). Therefore, it is imperative that complexation of organic chemicals with DOM be modelled correctly. In particular, contradictory research results can be reconciled by considering that DOM is not homogeneous. For instance, refractory humic acids, derived from decomposition of terrestrial and wetland organic material, are quite different from labile exudates from algae and other indigenous organisms.

Humic acids exhibit high polarity and do not readily complex neutral compounds. Natural humic acids from a Finnish lake with extensive marshes were spiked with a PCB, but a PCB-humic acid complex could not be demonstrated (Maaret et al. 1992). In another study, Freidig et al. (1998) used artificially prepared Aldrich humic acid to determine a humic acid-DOC partition coefficient (n = 5, n = 0.80), although they cautioned about extrapolation to the field:

$$KOC_{RefrDOM} = 28.84 \cdot KOW^{0.67} \tag{217}$$

where:

 $KOC_{RefrDOM}$ = refractory dissolved organic carbon partition coefficient (L/kg).

Until a better relationship is found, we are using a generalization of their equation to include polar compounds, transformed from organic carbon to organic matter, in AQUATOX:

$$KOM_{RefrDOM} = (28.84 \cdot KOW^{0.67} \cdot Nondissoc + (1 - Nondissoc) \cdot IonCorr \cdot 28.84 \cdot KOW^{0.67}) \cdot 0.526$$
(218)

where:

 $KOM_{RefrDOM}$ = refractory dissolved organic matter partition coefficient (L/kg).

Nonpolar lipids in algae occur in the cell contents, and it is likely that they constitute part of the labile dissolved exudate, which may be both excreted and lysed material. Therefore, the stronger relationship reported by Koelmans and Heugens (1998) for partitioning to algal exudate (n = 6, $r^2 = 0.926$) is:

$$KOC_{LabDOC} = 0.88 \cdot KOW$$
 (219)

which we also generalized for polar compounds and transformed:

$$KOM_{LabDOM} = (0.88 \cdot KOW \cdot Nondissoc + (1 - Nondissoc) \cdot IonCorr \cdot 0.88 \cdot KOW) \cdot 0.526$$
(220)

where:

 KOC_{LabDOC} = partition coefficient for labile dissolved organic carbon (L/kg); and KOM_{LabDOM} = partition coefficient for labile dissolved organic matter (L/kg).

Unfortunately, older data and modeling efforts failed to distinguish between hydrophobic compounds that were truly dissolved and those that were complexed with DOM. For example, the PCB water concentrations for Lake Ontario, reported by Oliver and Niimi (1988) and used by many subsequent researchers, included both dissolved and DOC-complexed PCBs (a fact which they

recognized). In their steady-state model of PCBs in the Great Lakes, Thomann and Mueller (1983) defined "dissolved" as that which is not particulate (passing a 0.45 micron filter). In their Hudson River PCB model, Thomann et al. (1991) again used an operational definition of dissolved PCBs. AQUATOX distinguishes between truly dissolved and complexed compounds; therefore, the partition coefficients calculated by AQUATOX may be larger than those used in older studies.

Bioaccumulation of PCBs in algae depends on solubility, hydrophobicity and molecular configuration of the compound, and growth rate, surface area and type, and content and type of lipid in the alga (Stange and Swackhamer 1994). Phytoplankton may double or triple in one day and periphyton turnover may be so rapid that some PCBs will not reach equilibrium (cf. Hill and Napolitano 1997); therefore, one should use the term "bioaccumulation factor" (BAF) rather than "bioconcentration factor," which implies equilibrium (Stange and Swackhamer 1994).

Hydrophobic compounds partition to lipids in algae, but the relationship is not a simple one. Phytoplankton lipids can range from 3 to 30% by weight (Swackhamer and Skoglund 1991), and not all lipids are the same. Polar phospholipids occur on the surface. Hydrophobic compounds preferentially partition to internal neutral lipids, but those are usually a minor fraction of the total lipids, and they vary depending on growth conditions and species (Stange and Swackhamer 1994). Algal lipids have a much stronger affinity for hydrophobic compounds than does octanol, so that the algal $BAF_{lipid} > K_{OW}$ (Stange and Swackhamer 1994, Koelmans et al. 1995, Sijm et al. 1998).

For algae, the approximation to estimate the dry-weight bioaccumulation factor ($r^2 = 0.87$), computed from Swackhamer and Skoglund's (1993) study of numerous PCB congeners, is:

$$\log(BCF_{Alga}) = 0.41 + 0.91 \cdot LogKOW \tag{221}$$

where:

$$BCF_{Alga}$$
 = partition coefficient between algae and water (L/kg).

Rearranging and extending to hydrophilic and ionized compounds:

$$BCF_{Alga} = 2.57 \cdot KOW^{0.91} \cdot Nondissoc + (1 - Nondissoc) \cdot IonCorr \cdot 2.57 \cdot KOW^{0.91}$$
 (222)

Comparing the results of using these coefficients, we see that they are consistent with the relative importance of the various substrates in binding organic chemicals (**Figure 74**). Binding capacity of detritus is greater than dissolved organic matter in Great Lakes waters (Stange and Swackhamer 1994, Gilek et al. 1996). In a study using Baltic Sea water, less than 7% PCBs were associated with dissolved organic matter and most were associated with algae (Björk and Gilek 1999). In contrast, in a study using algal exudate and a PCB, 98% of the dissolved concentration was as a dissolved organic matter complex and only 2% was bioavailable (Koelmans and Heugens 1998).

The influence of substrate polarity is evident in **Figure 75**, which shows the effect of ionization on binding of pentachlorophenol to various types of organic matter. The polar substrates, such as algal detritus, have an inflection point which is one pH unit higher than that of nonpolar substrates, such as refractory detritus. The relative importance of the substrates for binding is also demonstrated quite clearly.

For macrophytes, an empirical relationship reported by Gobas et al. (1991) for 9 chemicals with LogKOWs of 4 to 8.3 ($r^2 = 0.97$) is used:

$$\log(KB_{Macro}) = 0.98 \cdot LogKOW - 2.24 \tag{223}$$

Again, rearranging and extending to hydrophilic and ionized compounds:

$$KB_{Macro} = 0.00575 \cdot KOW^{0.98} \cdot (Nondissoc + 0.2)$$

For the invertebrate bioconcentration factor, the following empirical equation is used, based on 7 chemicals with LogKOWs ranging from 3.3 to 6.2 and bioconcentration factors for Daphnia pulex ($r^2 = 0.85$; Southworth et al., 1978; see also Lyman et al., 1982), converted to dry weight:

$$log(KB_{Invertebrate}) = (0.7520 \cdot LogKOW - 0.4362) \cdot WetToDry$$
 (225)

where:

 $KB_{Invertebrate}$ = partition coefficient between invertebrates and water (L/kg); and WetToDry = wet to dry conversion factor (unitless, default = 5).

Extending and generalizing to ionized compounds:

$$KB_{Invertebrate} = 0.3663 \cdot KOW^{0.7520} \cdot (Nondissoc + 0.01)$$

Figure 74
Partitioning to Various Types of Organic
Matter as a Function of KOW

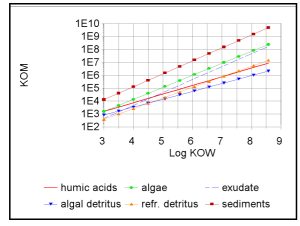
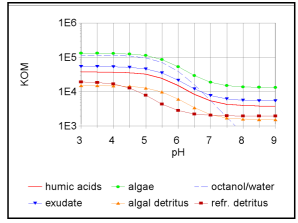


Figure 75
Partitioning to Various Types of Organic
Matter as a Function of pH



Fish take longer to reach equilibrium with the surrounding water; therefore, a nonequilibrium bioconcentration factor is used. For each pollutant, a whole-fish bioconcentration factor is based on the lipid content of the fish extended to hydrophilic chemicals (McCarty et al., 1992), with provision for ionization:

$$KB_{Fish} = Lipid \cdot WetToDry \cdot KOW \cdot (Nondissoc + 0.01)$$
 (227)

where:

 KB_{Fish} = partition coefficient between whole fish and water (L/kg);

Lipid = fraction of fish that is lipid (g lipid/g fish); and WetToDry = wet to dry conversion factor (unitless, default = 5).

Lipid content of fish is varied depending on the potential for growth as predicted by the bioenergetics equations; the initial lipid values for the species are given. The bioconcentration factor is adjusted for the time to reach equilibrium as a function of the clearance or elimination rate and the time of exposure (Hawker and Connell, 1985; Connell and Hawker, 1988; **Figure 76**):

$$BCF_{Fish} = KB_{Fish} \cdot (1 - e^{(-Depuration \cdot TElapsed)})$$
 (228)

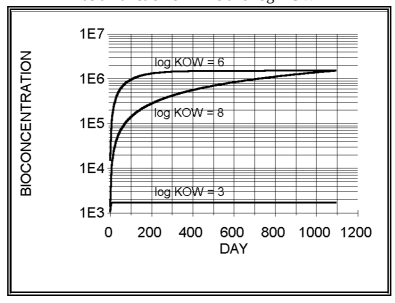
where:

 BCF_{Fish} = quasi-equilibrium bioconcentration factor for fish (L/kg);

TElapsed = time elapsed since fish was first exposed (d); and

Depuration = clearance, which may include biotransformation, see (258) (1/d).

Figure 76
Bioconcentration Factor for Fish as a Function of Time and log KOW



The concentration in each carrier is given by:

$$PPB_{i} = \frac{ToxState_{i}}{CarrierState_{i}} \cdot 1e6$$
 (229)

where:

 PPB_i = concentration of chemical in carrier i (µg/kg);

 $ToxState_i$ = mass of chemical in carrier i (ug/L); CarrierState = biomass of carrier (mg/L); and 1e6 = conversion factor (mg/kg).

7.7 Nonequilibrium Kinetics

Often there is an absence of equilibrium due to growth or insufficient exposure time, metabolic biotransformation, dietary exposure, and nonlinear relationships for very large and/or superhydrophobic compounds (Bertelsen et al. 1998). Although it is important to have a knowledge of equilibrium partitioning because it is an indication of the condition toward which systems tend (Bertelsen et al. 1998), it is often impossible to determine steady-state potential due to changes in bioavailability and physiology (Landrum 1998). For example, PCBs may not be at steady state even in large systems such as Lake Ontario that have been polluted over a long period of time. In fact, PCBs in Lake Ontario exhibit a 25-fold disequilibrium (Cook and Burkhard 1998). The challenge is to obtain sufficient data for a kinetic model (Gobas et al. 1995).

Sorption and Desorption to Sedimented Detritus

Partitioning to sediments appears to involve rapid sorption to particle surfaces, followed by slow movement into, and out of, organic matter and porous aggregates (Karickhoff and Morris, 1985). Therefore attainment of equilibrium may be slow. This applies to suspended detritus compartments as well. Because of the need to represent sorption and desorption separately in detritus, kinetic formulations are used (Thomann and Mueller, 1987), with provision for ionization:

Sorption =
$$kI_{Detr} \cdot Toxicant_{Water} \cdot DiffI_{Carrier} \cdot (Nondissoc + 0.01)$$

 $\cdot Org2C \cdot Detr \cdot 1e - 6$ (230)

$$Desorption = k2_{Detr} \cdot Diff2_{Carrier} \cdot Toxicant_{Detr}$$
 (231)

where:

Sorption = rate of sorption to given detritus compartment (μ g/L·d);

 kI_{Detr} = sorption rate constant (L/kg·d);

Nondissoc = fraction not ionized (unitless);

 $Toxicant_{Water}$ = concentration of toxicant in water ($\mu g/L$);

 $Diffl_{Carrier}$ = factor to normalize rate constant for given carrier (detritus compartment

in this case) based on all competing uptake rates (unitless);

 $Diff2_{Carrier}$ = factor to normalize loss rates (unitless);

Org2C = conversion factor for organic matter to carbon (= 0.526 g C/g organic

matter);

Detr = mass of each of the detritus compartments per unit volume (mg/L);

1e -6 = units conversion (kg/mg);

Desorption = rate of desorption from given sediment detritus compartment ($\mu g/L \cdot d$);

 $k2_{Detr}$ = desorption rate constant (1/d); and

 $Toxicant_{Detr}$ = mass of toxicant in each of the detritus compartments ($\mu g/L$).

Because there are several processes competing for the dissolved toxicant, the rate constants for these processes are normalized in order to preserve mass balance. The *Diff1* factor is computed for each direct uptake process, including sorption to detritus and algae, uptake by macrophytes, and uptake across animals' gills:

$$Diffl_{Carrier} = \frac{RateDiff_{Carrier}}{\sum RateDiff_{Carrier}}$$
(232)

$$RateDiff_{Carrier} = Gradient1_{Carrier} \cdot k1$$
 (233)

$$Gradient1_{Carrier} = \frac{Toxicant_{Water} \cdot kp_{Carrier} - PPB_{Carrier}}{Toxicant_{Water} \cdot kp_{Carrier}}$$
(234)

where:

 $RateDiff1_{Carrier}$ = maximum rate constant for uptake given the concentration gradient

 $(L/kg\cdot d);$

Gradient 1_{Carrier} = gradient between potential and actual concentrations of toxicant in

each carrier (unitless);

 $kp_{Carrier}$ = partition coefficient or bioconcentration factor for each carrier (L/kg);

 $PPB_{Carrier}$ = concentration of toxicant in each carrier (µg/kg).

Likewise, the loss rate constants are normalized; the equations parallel those for uptake, with the gradient being reversed:

$$Diff2_{Carrier} = \frac{RateDiff2_{Carrier}}{\sum RateDiff2_{Carrier}}$$
(235)

$$RateDiff2_{Carrier} = Gradient2_{Carrier} \cdot k2$$
 (236)

$$Gradient2_{Carrier} = \frac{PPB_{Carrier} - (PPB_{Water} \cdot kp_{Carrier})}{PPB_{Carrier}}$$
(237)

where:

 $RateDiff2_{Carrier} = maximum rate constant for loss given the concentration gradient$

(L/kg·d); and

Gradient2_{Carrier} = gradient between actual and potential concentrations of toxicant in

each carrier (unitless).

Desorption of the nonlabile or slow compartment is the reciprocal of the reaction time, which Karickhoff and Morris (1985) found to be a linear function of the partition coefficient over three orders of magnitude ($r^2 = 0.87$):

$$\frac{1}{k2} \approx 0.03 \cdot 24 \cdot KPSed \tag{238}$$

So *k2* is taken to be:

$$k2 = \frac{1.39}{KPSed} \tag{239}$$

where:

KPSed = detritus-water partition coefficient (L/kg, see **Eq. (212)**); and

= conversion from hours to days.

The nonlabile compartment may be involved in 40 to 90% of the sorption so, as a simplification, fast desorption of the labile compartment is ignored. This compensates in part for the fact that AQUATOX models the top layer of bottom sediments as if it were in close contact with the overlying water column (interstitial water is not modeled at this time).

The sorption rate constant k1 is set to 1200 L/kg*d, representing the very fast labile sorption of most chemicals.

Bioconcentration in Macrophytes and Algae

Macrophytes—As Gobas et al. (1991) have shown, submerged aquatic macrophytes take up and release organic chemicals over a measurable period of time at rates related to the octanol-water partition coefficient. Uptake and elimination are modeled assuming that the chemical is transported through both aqueous and lipid phases in the plant, with rate constants using empirical equations fit to observed data (Gobas et al., 1991), modified to account for ionization effects (**Figure 77**, **Figure 78**):

$$MacroUptake = k1 \cdot Diffl_{Plant} \cdot Toxicant_{Water} \cdot StVar_{Plant} \cdot 1e - 6$$
 (240)

$$Depuration_{Plant} = k2 \cdot Toxicant_{Plant} \cdot Diff2_{Plant}$$
 (241)

$$k1 = \frac{1}{0.0020 + \frac{500}{KOW \cdot Nondissoc}} \tag{242}$$

$$k2 = \frac{1}{1.58 + 0.000015 \cdot KOW \cdot Nondissoc}$$
 (243)

where:

MacroUptake = uptake of toxicant by plant ($\mu g/L \cdot d$);

Depuration_{Plant} = clearance of toxicant from plant (μ g/L·d);

 $StVar_{Plant}$ = biomass of given plant (mg/L); 1 e -6 = units conversion (kg/mg);

 $Toxicant_{Plant}$ = mass of toxicant in plant (μ g/L); kl = sorption rate constant (L/kg·d); k2 = elimination rate constant (1/d);

 $Diffl_{Plant}$ = factor to normalize uptake rates (unitless); $Diffl_{Plant}$ = factor to normalize loss rates (unitless);

KOW = octanol-water partition coefficient (unitless); and

Nondissoc = fraction of un-ionized toxicant (unitless).

Figure 77
Uptake Rate Constant for Macrophytes
(after Gobas et al., 1991)

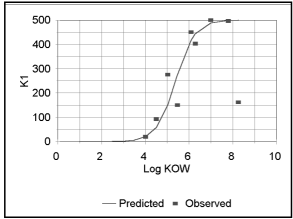
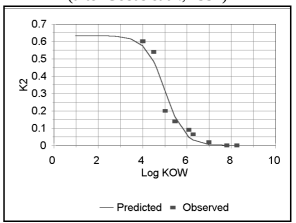


Figure 78
Elimination Rate Constant for Macrophytes
(after Gobas et al., 1991)



Algae—There is probably a two-step algal bioaccumulation mechanism for hydrophobic compounds, with rapid surface sorption of 40-90% within 24 hours and then a small, steady increase with transfer to interior lipids for the duration of the exposure (Swackhamer and Skoglund 1991). Uptake increases with increase in the surface area of algae (Wang et al. 1997). Therefore, the smaller the organism the larger the uptake rate constant (Sijm et al. 1998). However, in small phytoplankton, such as the nannoplankton that dominate the Great lakes, a high surface to volume ratio can increase sorption, but high growth rates can limit internal contaminant concentrations (Swackhamer and Skoglund 1991). The combination of lipid content, surface area, and growth rate results in species differences in bioaccumulation factors among algae (Wood et al. 1997). Uptake of toxicants is a function of the uptake rate constant and the concentration of toxicant truly dissolved in the water, and constrained by competitive uptake by other compartments:

$$AlgalUptake = k1 \cdot Michaelis \cdot Diff1 \cdot ToxState \cdot Carrier \cdot 1e - 6$$
 (244)

where:

AlgalUptake = rate of sorption by algae (μ g/L-d);

kl = uptake rate constant (L/kg-d), see (245);

Michaelis = Michaelis-Menton construct for nonlinear uptake (unitless), see (246);

Diff1 = factor to normalize uptake rates (unitless), see (232);

 $ToxState = concentration of dissolved toxicant (<math>\mu g/L$); Carrier = biomass of algal compartment (mg/L); and

1e-6 = conversion factor (kg/mg).

The kinetics of partitioning of toxicants to algae is based on studies on PCB congeners in The Netherlands by Koelmans, Sijm, and colleagues and at the University of Minnesota by Skoglund and Swackhamer. Both groups found uptake to be very rapid. Sijm et al. (1998) presented data on several congeners that were used in this study to develop the following relationship for phytoplankton (**Figure 79**):

$$kI = \frac{1}{1.8E-6 + 1/(KOW \cdot Nondissoc)}$$
 (245)

Because size-dependent passive transport is indicated (Sijm et al., 1998), uptake by periphyton is set arbitrarily at ten percent of that for phytoplankton.

In order to represent saturation kinetics, *Michaelis* is computed as:

$$Michaelis = \frac{BCF_{Algae} \cdot ToxState - PPB_{Algae}}{BCF_{Algae} \cdot ToxState}$$
(246)

where:

 BCF_{Algae} = steady-state bioconcentration factor for algae (L/kg), see (222); and PPB_{Algae} = concentration of toxicant in algae (mg/kg).

Depuration is modeled as a linear function; it does not include loss due to excretion of photosynthate with associated toxicant, which is modeled separately.

$$Depuration = k2 \cdot State \tag{247}$$

where:

Depuration = elimination of toxicant ($\mu g/L-d$);

State = concentration of toxicant associated with alga ($\mu g/L$); and

k2 = elimination rate constant (1/d).

Based in part on Skoglund et al. (1996), but ignoring surface sorption and recognizing that growth dilution is explicit in AQUATOX, the elimination rate constant (**Figure 80**) is computed as:

$$k2 = \frac{k1}{KOW} \tag{248}$$

Aside from obvious structural differences, algae may have very high lipid content (20% for *Chlorella* sp. according to Jørgensen et al., 1979) and macrophytes have a very low lipid content (0.2% in *Myriophyllum spicatum* as observed by Gobas et al. 1991), which affect both uptake and elimination of toxicants.

Figure 79
Algal Sorption Rate Constant as a Function of Octanol-water Partition Coefficient

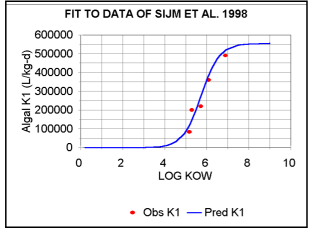
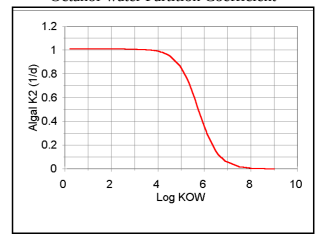


Figure 80
Rate of Elimination by Algae as a Function of Octanol-water Partition Coefficient



Bioaccumulation in Animals

Animals can absorb toxic organic chemicals directly from the water through their gills and from contaminated food through their guts. Direct sorption onto the body is ignored as a simplifying assumption in this version of the model. Reduction of body burdens of organic chemicals is accomplished through excretion and biotransformation, which are often considered together as empirically determined elimination rates. "Growth dilution" occurs when growth of the organism is faster than accumulation of the toxicant. Gobas (1993) includes fecal egestion, but in AQUATOX egestion is merely the amount ingested but not assimilated; it is accounted for indirectly in *DietUptake*. However, fecal loss is important as an input to the detrital toxicant pool, and it is considered later in that context. Inclusion of mortality and promotion terms is necessary for mass balance, but emphasizes the fact that average concentrations are being modeled for any particular compartment.

Gill Sorption—An important route of exposure is by active transport through the gills (Macek et al., 1977). This is the route that has been measured so often in bioconcentration experiments with fish. As the organism respires, water is passed over the outer surface of the gill and blood is moved past the inner surface. The exchange of toxicant through the gill membrane is assumed to be facilitated by the same mechanism as the uptake of oxygen, following the approach of Fagerström and Åsell (1973, 1975), Weininger (1978), and Thomann and Mueller (1987; see also Thomann, 1989). Therefore, the uptake rate for each animal can be calculated as a function of respiration (Leung, 1978; Park et al., 1980):

$$GillUptake = KUptake \cdot Toxicant_{Water} \cdot Diffl_{Carrier}$$
 (249)

$$KUptake = \frac{WEffTox \cdot Respiration \cdot O2Biomass}{Oxygen \cdot WEffO2}$$
(250)

where:

 $GillUptake = uptake of toxicant by gills (\mu g/L - d);$

KUptake = uptake rate (1/d);

 $Toxicant_{Water}$ = concentration of toxicant in water ($\mu g/L$);

 $Diffl_{Carrier}$ = factor to normalize rate constant for given carrier (animal compartment

in this case) based on all competing uptake rates (unitless), see (232);

WEffTox = withdrawal efficiency for toxicant by gills (unitless), see (251);

Respiration = respiration rate (mg biomass/L·d), see (76);

O2Biomass = ratio of oxygen to organic matter (mg oxygen/mg biomass; generally

0.575);

Oxygen = concentration of dissolved oxygen (mg oxygen/L); and WEffO2 = withdrawal efficiency for oxygen (unitless, generally 0.62).

The oxygen uptake efficiency *WEffO2* is assigned a constant value of 0.62 based on observations of McKim et al. (1985). The toxicant uptake efficiency, *WEffTox*, can be expected to have a sigmoidal relationship to the log octanol-water partition coefficient based on aqueous and lipid transport (Spacie and Hamelink, 1982). This is represented by an inelegant but reasonable, piece-wise fit (**Figure 81**) to the data of McKim et al. (1985) using 750-g fish, corrected for ionization:

If LogKOW < 1.5 then

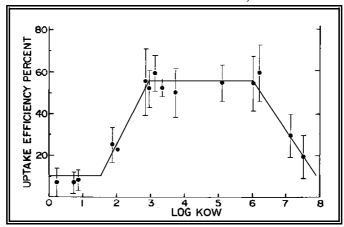
$$\overline{WEffTox} = 0.1$$
If $1.5 \le LogKOW > 3.0$ then
$$WEffTox = 0.1 + Nondissoc \cdot (0.3 \cdot LogKOW - 0.45)$$
If $3.0 \le LogKOW \le 6.0$ then
$$WEffTox = 0.1 + Nondissoc \cdot 0.45$$
If $6.0 < LogKOW < 8.0$ then
$$WEffTox = 0.1 + Nondissoc \cdot (0.45 - 0.23 \cdot (LogKOW - 6.0))$$
(251)

If $LogKOW \ge 8.0$ then WEffTox = 0.1

where:

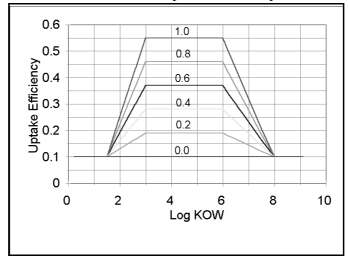
LogKOW = log octanol-water partition coefficient (unitless); and Nondissoc = fraction of toxicant that is un-ionized (unitless), see (188).

Figure 81
Piece-wise Fit to Observed Toxicant Uptake Data;
Modified from McKim et al., 1985



Ionization decreases the uptake efficiency (**Figure 82**). This same algorithm is used for invertebrates. Thomann (1989) has proposed a similar construct for these same data and a slightly different construct for small organisms, but the scatter in the data do not seem to justify using two different constructs.

Figure 82
The Effect of Differing Fractions of Un-ionized
Chemical on Uptake Efficiency



Dietary Uptake—Hydrophobic chemicals usually bioaccumulate primarily through absorption from contaminated food. Persistent, highly hydrophobic chemicals demonstrate biomagnification or increasing concentrations as they are passed up the food chain from one trophic level to another; therefore, dietary exposure can be quite important (Gobas et al., 1993). Uptake from contaminated prey can be computed as (Thomann and Mueller, 1987; Gobas, 1993):

$$DietUptake_{Prey} = KD_{Prey} \cdot PPB_{Prey} \cdot 1e - 6$$
 (252)

$$KD_{Prey} = GutEffTox \cdot Ingestion_{Prey}$$
 (253)

where:

 $DietUptake_{Prey}$ = uptake of toxicant from given prey (μ g toxicant/L·d); KD_{Prey} = dietary uptake rate for given prey (mg prey/L·d);

 PPB_{Prev} = concentration of toxicant in given prey (μ g toxicant/kg prey);

1 e-6 = units conversion (kg/mg);

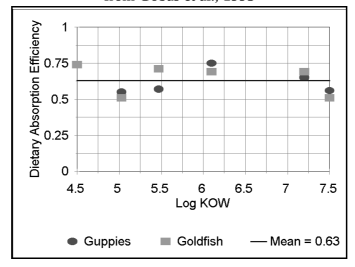
GutEffTox = efficiency of sorption of toxicant from gut (unitless); and

 $Ingestion_{Prev}$ = ingestion of given prey (mg prey/L·d), see (68).

Gobas (1993) presents an empirical equation for estimating *GutEffTox* as a function of the octanol-water partition coefficient. However, data published by Gobas et al. (1993) suggest that there is no trend in efficiency between *LogKOW* 4.5 and 7.5 (**Figure 83**); this is to be expected because the digestive system has evolved to assimilate a wide variety of organic molecules. Therefore, the mean value of 0.63 is used in AQUATOX as a constant for small fish. Nichols et al. (1998) demonstrated that uptake is more efficient in larger fish; therefore, a value of 0.90 is used for large game fish because of their size. Invertebrates generally exhibit lower efficiencies; Landrum and Robbins (1990) showed that values ranged from 0.42 to 0.24 for chemicals with log KOWs from 4.4 to 6.7; the mean value of 0.35 is used for invertebrates in AQUATOX.

Figure 83

GutEffTox Constant Based on Mean Value for Data from Gobas et al., 1993



Elimination—Elimination or clearance includes both excretion (depuration) and biotransformation of a toxicant by organisms. Biotransformation may cause underestimation of elimination (McCarty et al., 1992). Therefore, an overall elimination rate constant is estimated and reported in the toxicity record. The user may then modify the value based on observed data; that value is used in subsequent simulations. However, because biotransformation is not modeled explicitly, this is treated as depuration for purposes of tracking the transfer of toxicant.

For purposes of estimating elimination, a modification of **Eq. (250)** is used to compute uptake, assuming an allometric relationship between respiration and the weight of the animal (Thomann, 1989):

$$k1 = 1000 \cdot WetWt^{-0.2} \cdot WEffTox \tag{254}$$

where:

k1 = uptake rate constant (L/kg·d); WetWt = mean wet weight of organism (g);

1000 = units conversion (g/kg);

WEffTox = withdrawal efficiency for toxicant by gills, see Eq. (251) (unitless).

If, as Thomann (1989) assumes, lipid-normalized bioconcentration is equal to the octanol-water partition coefficient at equilibrium and zero growth, then:

$$k2 = \frac{k1}{KOW \cdot LipidFrac \cdot WetToDry \cdot (Nondissoc + 0.1)}$$
 (255)

where:

k2 = elimination rate constant (1/d); and

KOW = octanol-water partition coefficient (unitless);
LipidFrac = fraction of lipid in organism (g lipid/g organism);

Wet2Dry = wet to dry weight ratio (5); and

Nondissoc = fraction of compound un-ionized (unitless).

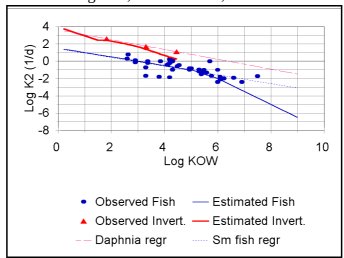
This simple relationship, although weak, has been used in AQUATOX for both invertebrates and fish (**Figure 84**). However, the fish curve seems to drastically underestimate clearance at higher KOWs. Therefore, as an alternative until the formulation is changed, K2 estimates may be entered manually using as guides regression equations for *Daphnia*:

$$\text{Log } k2 = -0.5688 \cdot \text{Log } KOW + 3.6445$$

and small fish:

$$Log k2 = -0.503 \cdot Log KOW + 1.45$$

Figure 84
Depuration Rate Constants for *Daphnia* and for 10-g Fish; see Thomann, 1989



For any given time the clearance rate is:

$$Depuration_{Animal} = k2 \cdot Toxicant_{Animal}$$
 (258)

where:

 $Depuration_{Animal}$ = clearance rate ($\mu g/L \cdot d$); and

 $Toxicant_{Animal}$ = mass of toxicant in given animal ($\mu g/L$).

Linkages to Detrital Compartments

Toxicants are transferred from organismal to detrital compartments through defecation and mortality. The amount transferred due to defecation is the unassimilated portion of the toxicant that is ingested:

$$DefecationTox = \sum (KEgest_{Pred, Prey} \cdot PPB_{Prey} \cdot 1e - 6)$$
 (259)

$$KEgest_{Pred, Prey} = (1 - GutEffTox) \cdot Ingestion_{Pred, Prey}$$
 (260)

where:

DefecationTox = rate of transfer of toxicant due to defecation ($\mu g/L \cdot d$);

 $KEgest_{Pred, Prey}$ = fecal egestion rate for given prey by given predator (mg

prey/L·d);

 PPB_{Prev} = concentration of toxicant in given prey ($\mu g/kg$);

1 e-6 = units conversion (kg/mg);

GutEffTox = efficiency of sorption of toxicant from gut (unitless); and rate of ingestion of given prey by given predator (mg/L·d), see

(68).

The amount of toxicant transferred due to mortality may be large; it is a function of the concentrations of toxicant in the dying organisms and the mortality rates:

$$MortTox = \sum (Mortality_{Org} \cdot PPB_{Org} \cdot 1e-6)$$
 (261)

where:

MortTox = rate of transfer of toxicant due to mortality (μ g/L·d); $Mortality_{Org}$ = rate of mortality of given organism (mg/L·d), see(80); PPB_{Org} = concentration of toxicant in given organism (μ g/kg); and

1 e-6 = units conversion (kg/mg).

8. ECOTOXICOLOGY

8.1 Acute Toxicity of Compounds

Toxicity is based on the internal concentration of the toxicant in the specified organism. Many compounds, especially those with higher octanol-water partition coefficients, take appreciable time to accumulate in the tissue. Therefore, length of exposure is critical in determining toxicity. Although AQUATOX cannot currently model mercury, mercury is used as an example in the following discussion because of the availability of excellent data. The same principles apply to organic toxicants and to both plants and animals.

Based on the Mancini (1983) model, the lethal internal concentration of a toxicant can be expressed as (Crommentuijn et al. (1994):

$$LethalConc = \frac{LCInfinite}{1 - e^{-k2 \cdot TElapsed}}$$
 (262)

where:

LethalConc = tissue-based concentration of toxicant that causes mortality (ppb

or $\mu g/kg$);

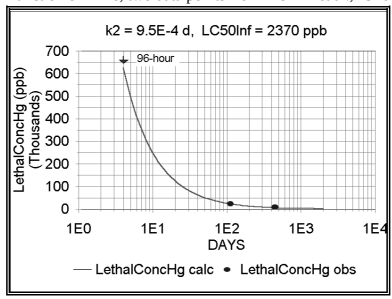
LCInfinite = ultimate internal lethal toxicant concentration after an infinitely

long exposure time (ppb);

k2 = elimination rate constant (1/d); and

TElapsed = period of exposure (d).

Figure 85
Lethal Concentration of MeHg in Brook Trout as a
Function of Time; two data points from McKim et al., 1976



The model estimates k2, see (242), (248), and (255), assuming that this k2 is the same as that measured in bioconcentration tests; good agreement has been reported between the two (Mackay et al., 1992). The user may then override that estimate by entering an observed value. The k2 can be calculated based on the observed half-life:

$$k2 = \frac{0.693}{t_{1/2}} \tag{263}$$

where:

 t_{ν_2} = observed half-life.

Exposure is limited to the lifetime of the organism:

If
$$TElapsed > LifeSpan$$
 Then $TElapsed = LifeSpan$ (264)

where:

LifeSpan = user-defined mean lifetime for given organism (d).

Based on an estimate of time to reach equilibrium (Connell and Hawker, 1988),

if
$$TElapsed > \frac{4.605}{k2}$$
 then
$$LethalConc = LCInfinite$$
(265)

LCInfinite can be computed by a rearrangement of Eq. (262).

$$LCInfinite = LethalConc \cdot (1 - e^{-k2 \cdot ObsTElapsed})$$
 (266)

where:

ObsTElapsed = exposure time in toxicity determination (h).

The internal lethal concentration for a given period of exposure can be computed from reported acute toxicity data based on the simple relationship suggested by an algorithm in the FGETS model (Suárez and Barber, 1992):

$$LethalConc = BCF \cdot LC50 \tag{267}$$

where:

BCF = bioconcentration factor (L/kg), see (222), (228); and LC50 = concentration of toxicant in water that causes 50% mortality (μ g/L).

A given *LC50* can be provided by the user, or the user may choose to have the model estimate the *LC50* based on regressions from other species for which there are data (Mayer and Ellersieck, 1986, Suter et al., 1986).

The fraction killed by a given internal concentration of toxicant and the time-dependent *LCInfinite* is best estimated using the cumulative form of the Weibull distribution (Mackay et al., 1992):

$$CumFracKilled = 1 - e^{-\left(\frac{PPB}{LethalConc}\right)^{1/Shape}}$$
(268)

where:

CumFracKilled = cumulative fraction of organisms killed,

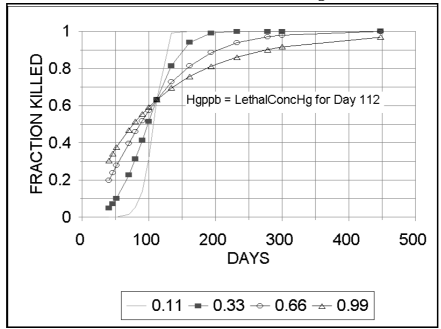
PPB = internal concentration of toxicant ($\mu g/kg$); and

Shape = parameter expressing variability in toxic response (unitless).

As a practical matter, if *CumFracKilled* exceeds 95%, then it is set to 100% to avoid complex computations with small numbers. By setting organismal loadings to very small numbers, seed values can be maintained in the simulation.

The *Shape* parameter is important because it controls the spread of mortality. The larger the value, the greater the distribution of mortality over toxicant concentrations and time. Methyl mercury toxicity exhibits a rapid response over a short time period, so *Shape* has a value of about 0.1 (**Figure 86**). However, Mackay et al. (1992) found that a value of 0.33 gave the best fit to data on toxicity of 21 narcotic chemicals to fathead minnows. This value is used as a default in AQUATOX, but it can be changed by the user. Note that when the internal concentration equals the lethal concentration the Weibull equation predicts 63% mortality.

Figure 86
The Effect of *Shape* on the Variability of Response to a Given Internal Concentration of MeHg



The loss due to acute mortality accounts for the population already exposed and subject only to increased exposure, and the newly exposed population that is more vulnerable. Mortality is computed as:

$$Poisoned = Resistant \cdot FracKilled + Nonresistant \cdot CumFracKilled$$
 (269)

where:

Poisoned = biomass of given organisms killed by exposure to toxicant at given time

(mg/L);

Resistant = biomass not killed by previous exposure (mg/L);

FracKilled = fraction killed in excess of the previous fraction (unitless); and

Nonresistant = biomass not previously exposed; the biomass in excess of the resistant

biomass (mg/L).

8.2 Chronic Toxicity

Organisms usually have adverse reactions to toxicants at levels significantly below those that cause death. In fact, the acute to chronic ratio is commonly used to quantify this relationship. Application factors (AFs), which are the inverse of the acute to chronic ratio, are employed in the model to predict chronic effect responses. The user supplies observed EC50 values, which are used to compute AFs. For example:

$$AFGrowth = \frac{EC50Growth}{LC50}$$
 (270)

where:

EC50Growth = external concentration of toxicant at which there is a 50% reduction in

growth (μ g/L);

AFGrowth = chronic to acute ratio for growth; and

LC50 = external concentration of toxicant at which 50% of population is killed

 $(\mu g/L)$.

If the user enters an observed EC50 value, the model provides the option of applying the resulting AF to estimate EC50s for other organisms. The computations for AFPhoto and AFRepro are similar.

Similar to computation of acute toxicity in the model, chronic toxicity is based on internal concentrations of a toxicant. Often chronic effects form a continuum with acute effects and the difference is merely one of degree (Mackay et al., 1992). Regardless of whether or not the mode of action is the same, the computed application factors relate the observed effect to the acute effect and permit efficient computation of chronic effects factors in conjunction with computation of acute effects. Because AQUATOX simulates biomass, no distinction is made between reduction in a process in an individual and the fraction of the population exhibiting that response. The commonly measured reduction in photosynthesis is a good example: the data only indicate that a given reduction takes place at a given concentration, not whether all individuals are affected. The application factor enters into the Weibull equation to estimate reduction factors for photosynthesis, growth, and reproduction:

$$FracPhoto = e^{-\left(\frac{PPB}{LethalConc \cdot AFPhoto}\right)^{1/Shape}}$$
 (271)

$$RedGrowth = 1 - e^{-\left(\frac{PPB}{LethalConc \cdot AFGrowth}\right)^{1/Shape}}$$
 (273)

$$RedRepro = 1 - e^{-\left(\frac{PPB}{LethalConc \cdot AFRepro}\right)^{1/Shape}}$$
 (272)

where:

FracPhoto = reduction factor for effect of toxicant on photosynthesis (unitless);

RedGrowth = factor for reduced growth in animals (unitless);

RedRepro = factor for reduced reproduction in animals (unitless);

PPB = internal concentration of toxicant ($\mu g/kg$);

LethalConc = tissue-based concentration of toxicant that causes mortality (μ g/kg); AFPhoto = chronic to acute ratio for photosynthesis (unitless, default of 0.10); AFGrowth = chronic to acute ratio for growth in animals (unitless, default of 0.10); chronic to acute ratio for reproduction in animals (unitless, default of

0.05); and

Shape = parameter expressing variability in toxic response (unitless, default of

0.33).

The reduction factor for photosynthesis, *FracPhoto*, enters into the photosynthesis equation (Eq. (26)), and it also appears in the equation for the acceleration of sinking of phytoplankton due to stress (Eq. (58)).

The variable for reduced growth, *RedGrowth*, is arbitrarily split between two processes, ingestion (Eq. **(68)**), where it reduces consumption by 20%:

$$ToxReduction = 1 - (0.2 \cdot RedGrowth)$$
 (274)

and egestion (Eq. (73)), where it increases the amount of food that is not assimilated by 80%:

$$IncrEgest = (1 - EgestCoeff_{prey, pred}) \cdot 0.8 \cdot RedGrowth$$
 (275)

These have indirect effects on the rest of the ecosystem through reduced predation and increased production of detritus in the form of feces.

Embryos are often more sensitive to toxicants, although reproductive failure may occur for various reasons. As a simplification, the factor for reduced reproduction, *RedRepro*, is used only to increase gamete mortality (Eq. (84)) beyond what would occur otherwise:

$$IncrMort = (1 - GMort) \cdot RedRepro$$
 (276)

By modeling chronic and acute effects, AQUATOX makes the link between chemical fate and the functioning of the aquatic ecosystem—a pioneering approach that has been refined over the past twelve years, following the first publications (Park et al., 1988; Park, 1990).

9. REFERENCES

- Abbott, J.D., S.W. Hinton, and D.L. Borton. 1995. Pilot Scale Validation of the RIVER/FISH Bioaccumulation Modeling Program for Nonpolar Hydrophobic Organic Compounds Using the Model Compounds 2,3,7,8-TCDD and 2,3,7,8-TCDF. *Environmental Toxicology and Chemistry*, 14(11):1999-2012.
- Alexander, M. 1977. *Introduction to Soil Microbiology, 2nd Edition*, Wiley & Sons, New York, 467 pp.
- Ambrose, R.B., Jr., T.A. Wool, J.L. Martin, J.P. Connolly, and R.W. Schanz. 1991. *WASP5.x, A Hydrodynamic and Water Quality Model—Model Theory, User's Manual, and Programmer's Guide*. U.S. Environmental Protection Agency, Environmental Research Laboratory, Athens, Georgia.
- Anonymous. 1988. *Latin.for*. Unpublished FORTRAN computer code, presumably developed at Oak Ridge National Laboratory.
- APHA (American Public Health Association). 1985. Standard Methods for the Examination of Water and Wastewater, 16th Edition, APHA, Washington D.C.
- Aruga, Y. 1965. Ecological Studies of Photosynthesis and Matter Production of Phytoplankton II. Photosynthesis of Algae in Relation to Light Intensity and Temperature. *Bot. Mag. Tokyo*, 78:360-365.
- Baca, R.G., and R.C. Arnett. 1976. *A Limnological Model for Eutrophic Lakes and Impoundments*. Batelle Pacific Northwest Laboratories, Richland, Washington.
- Barko, J.W., and R.M. Smart. 1980. Mobilization of sediment phosphorus by submersed freshwater macrophytes. *Freshwater Biology* 10:229-238.
- Berman, M.S., and S. Richman. 1974. The Feeding Behavior of *Daphnia pulex* from Lake Winnebago, Wisconsin. *Limnol. Oceanog.* 19:105-109.
- Bertelsen, S.L., A.D. Hoffman, C.A. Gallinat, C.M. Elonen, and J.W. Nichols. 1998. Evaluation of Log K_{OW} and Tissue Lipid Content as Predictors of Chemical Partitioning to Fish Tissues. *Environmental Toxicology and Chemistry*, 17(8):1447-1455.
- Bierman, V.J., Jr., D.M. Dolan, E.F. Stoermer, J.E. Gannon, and V.E. Smith. 1980. *The Development and Calibration of a Multi-Class Phytoplankton Model for Saginaw Bay, Lake Huron*. Great Lakes Environmental Planning Study, Contr. No. 33. Great Lakes Basin Commission, Ann Arbor, Michigan.

- Björk, M., and M Gilek. 1999. Efficiencies of Polychlorinated Biphenyl Assimilation from Water and Algal Food by the Blue Mussel (*Mytilus edulis*). *Environmental Toxicology and Chemistry*, 18(4):765-771.
- Bloomfield, J.A., R.A. Park, D. Scavia, and C.S. Zahorcak. 1973. Aquatic Modeling in the EDFB, US-IBP. in E.J. Middlebrooks, D.H. Falkenborg, and T.W. Maloney, Eds., *Modeling the Eutrophication Process*, Ann Arbor, Michigan: Ann Arbor Science Publishers.
- Bogdan, K.G., and D.C. McNaught. 1975. Selective Feeding by *Diaptomus* and *Daphnia. Verh. Intl. Ver. Limnol.* 19:2935-2942.
- Bowie, G.L., W.B. Mills, D.B. Porcella, C.L. Campbell, J.R. Pagenkopf, G.L. Rupp, K.M. Johnson, P.W.H. Chan and S.A. Gherini. 1985. *Rates, Constants, and Kinetics Formulations in Surface Water Quality Modeling (Ed. 2)*. U.S. Environmental Protection Agency, Athens, Georgia. EPA-600/3-85-040.
- Brandl, Z., and C.H. Fernando. 1975. Food Consumption and Utilization in Two Freshwater Cyclopoid Copepods (*Mesocyclops edax* and *Cyclops vicinus*). *Int. Rev. Ges. Hydrobiol*. 60:471-494.
- Bristow, J.M., and Mhitcombe. 1971. The Role of Roots in the Nutrition of Aquatic Vascular Plants. *Amer. Jour. Bot.*, 58:8-13.
- Brownlie, W.R. 1981. "Prediction of Flow Depth and Sediment Discharge in Open Channels". Report No. KH-R-43A. W.M. Keck laboratory of Hydraulics and Water Resources Division of Engineering and Applied Science. California Institute of Technology, Pasadena California.
- Burkhard, L.P. 1998. Comparison of Two Models for Predicting Bioaccumulation of Hydrophobic Organic Chemicals in a Great Lakes Food Web. *Environmental Toxicology and Chemistry*, 17(3):383-393.
- Burns, C.W. 1969. Particle Size and Sedimentation in the Feeding Behavior of Two Species of *Dahnia. Limnol. Oceanog.* 14:392-402.
- Burns, L.A., and D.M. Cline. 1985. *Exposure Analysis Modeling System, Reference Manual for EXAMS II*. U.S. Environmental Protection Agency, Athens, Georgia, EPA-600/3-85-038.
- Burns, L.A., D.M. Cline, and R.R. Lassiter. 1982. *Exposure Analysis Modeling System (EXAMS): User Manual and System Documentation*. U.S. Environmental Protection Agency, Athens, Georgia. EPA-600/3-82-023.

- Canale, R.P., L.M. Depalma, and A.H. Vogel. 1975. A Food Web Model for Lake Michigan. Part 2—Model Formulation and Preliminary Verification. Tech. Report 43, Michigan Sea Grant Program, MICU-SG-75-201.
- Canale, R.P., L.M. Depalma, and A.H. Vogel. 1976. A Plankton-Based Food Web Model for Lake Michigan, *in* R.P. Canale (ed.) *Modeling Biochemical Processes in Aquatic Ecosystems*. Ann Arbor Science Publishers, Ann Arbor, Michigan, pp. 33-74.
- Chapra, S.C., and K.H. Reckhow. 1983. *Engineering Approaches for Lake Management Volume 2: Mechanistic Modeling*. Butterworth Publishers, Boston, 492 pp.
- Chen, C.W. 1970. Concepts and Utilities of Ecologic Models. *Jour. San. Eng. Div. ASCE*, 96(SA 5):1085-1086.
- Chen, C.W., and G.T. Orlob. 1975. Ecological Simulation for Aquatic Environments, in B.C. Patton, Ed., *Systems Analysis and Simulation in Ecology Vol. III*, New York: Academic Press, pp. 476-588.
- Chow, V.T. 1959. Open Channel Hydraulics, McGraw-Hill, New York.
- Churchill, M.A.H.L. Elmore, and R.A. Buckingham. 1962. The Prediction of Stream Reaeration Rates, *ASCE*, *Journal Sanitary Engineering Division*, 88:SA4:1-46.
- Clesceri, L.S. 1980. Modeling the Fate of Synthetic Organic Chemicals, in A.W. Maki, K.L. Dickson, and J. Cairns, Jr. (eds.), *Biotransformation and Fate of Chemicals in the Aquatic Environment*, American Society for Microbiology, Washington, D.C.
- Clesceri L.S., R.A. Park, and J.A. Bloomfield. 1977. A General Model of Microbial Growth and Decomposition in Aquatic Ecosystems. *Applied and Environmental Microbiology* 33(5):1047–1058.
- Clifford, R., Jr. 1982. The Lake George Lay Monitoring Program 1981, In M.H. Schadler (ed.) *The Lake George Ecosystem Volume II*. The Lake George Association, Lake George, New York, pp. 126-130.
- Colby, J.A., and C.D. McIntire. 1978. *Mathematical Documentation for a Lotic Ecosystem Model*. Internal Report No. 165, Coniferous Forest Biome, Oregon State University, Corvallis, Oregon,
- Collins C.D. 1980. Formulation and Validation of a Mathematical Model of Phytoplankton Growth. *Ecology* 6:639-649.

- Collins, C.D., and R.A. Park. 1989. Chapter 15, Primary Productivity. In *Mathematical Submodels in Water Quality Systems*, S.E. Jørgensen and M.J. Gromiec, eds. Amsterdam: Elsevier, pp. 299-330.
- Collins, C.D., R.A. Park, and C.W. Boylen. 1985. *A Mathematical Model of Submersed Aquatic Plants*. Miscell. Paper A-85-2, U.S. Army Engineers Waterways Experiment Station, Vicksburg, Mississippi.
- Collins, C.D., and J.H. Wlosinski. 1983. *Coefficients for Use in the U.S. Army Corps of Engineers Reservoir Model, CE-QUAL-R1*. U.S. Army Engineer Waterways Experiment Station, Vicksburg, Mississippi, 120 pp.
- Connell, D.W., and D.W. Hawker. 1988. Use of Polynomial Expressions to Describe the Bioconcentration of Hydrophobic Chemicals by Fish. *Ecotoxicology and Environmental Safety* 16:242-257.
- Connolly, J.P., and D. Glaser. 1998. Use of Food Web Models to Evaluate Bioaccumulation. *National Sediment Bioaccumulation Conference Proceedings*. U.S. Environmental Protection Agency Office of Water EPA 823-R-98-002, p.4-5-4-17.
- Cook, P.M., and L.P. Burkhard. 1998. Development of Bioaccumulation Factors for Protection of Fish and Wildlife in the Great Lakes. *National Sediment Bioaccumulation Conference Proceedings*. U.S. Environmental Protection Agency Office of Water EPA 823-R-98-002, p. 3-19-3-27.
- Covar, A.P. 1976. Selecting the Proper Reaeration Coefficient for Use in Water Quality Models. Presented at US EPA Conference on Environmental Simulation and Modeling, April 19-22, Cincinnati, Ohio.
- Crommentuijn, T., C.J.A.M. Doodeman, A. Doornekamp, J.J.C. van der Pol, J.J.M. Bedaux, and C.A.M. van Gestel. 1994. Lethal Body Concentrations and Accumulation Patterns Determine Time-Dependent Toxicity of Cadmium in Soil Arthropods. *Environmental Toxicity and Chemistry* 13(11):1781-1789.
- DePinto, J.V. 1979. Water Column Death and Decomposition of Phytoplankton: An Experimental and Modeling Review. In *Pespectives on Lake Ecosystem Modeling*, D. Scavia and A. Robertson, eds. Ann Arbor, Mich.: Ann Arbor Science, pp. 25-52.
- Desormeau, C.J. 1978. *Mathematical Modeling of Phytoplankton Kinetics with Application to Two Alpine Lakes*. Report 4, Center for Ecological Modeling, 21 pp.
- Di Toro, D.M. 1985. A Particle Interaction Model of Reversible Organic Chemical Sorption. *Chemosphere*, 14(10):1503-1538.

- Di Toro, D.M., D.J. O'Connor, and R.V. Thomann. 1971. A Dynamic Model of the Phytoplankton Population in the Sacramento-San Joaquin Delta, in *Nonequilibrium Systems in Natural Water Chemistry*. Adv. Chem. Ser. 106, American Chemical Society, Washington, D.C., pp. 131-180.
- Effler, Steven W. 1996. *Limnological and Engineering Analysis of a Polluted Urban Lake*. New York: Springer-Verlag, 832 pp.
- Egglishaw, H.J. 1972. An Experimental Study of the Breakdown of Cellulose in Fast-Flowing Streams. In Melchiorri-Santolini, U., and J.W. Hopton, Eds., *Detritus and Its Role in Aquatic Ecosystems*, Proceedings of an IBP-UNESCO Symposium, *Mem. Ist. Ital. Idrobiol.* 29 Suppl.:405-428.
- Engelund, F. and E. Hansen. 1967. *A Monograph of Sediment Transport in Alluvial Streams*. Teknisk Vorlag, Copenhagen, Denmark.
- Environmental Laboratory. 1982. *CE-QUAL-R1: A Numerical One-Dimensional Model of Reservoir Water Quality; A User's Manual*. Instruction Report E-82-1, U.S. Army Engineers Waterways Experiment Station, Vicksburg, Miss.
- EPA. 1986. *Quality Criteria for Water, 1986*. Environmental Protection Agency, Washington, D.C., Office of Water Regulations and Standards, EPA/440/5-86/001, 398 pp.
- Fagerström, T., and B. Åsell. 1973. Methyl Mercury Accumulation in an Aquatic Food Chain, A Model and Some Implications for Research Planning. Ambio, 2(5):164-171.
- Fagerström, T., R. Kurtén ,and B. Åsell. 1975. Statistical Parameters as Criteria in Model Evaluation: Kinetics of Mercury Accumulation in Pike *Esox lucius*. Oikos 26:109-116.
- Fogg, G.E., C. Nalewajko, and W.D. Watt. 1965. Extracellular Products of Phytoplankton Photosynthesis. *Proc. Royal Soc. Biol.*, 162:517-534.
- Ford, D.E., and K.W. Thornton. 1979. Time and Length Scales for the One-Dimensional Assumption and Its Relation to Ecological Models. *Water Resources Research* 15(1):113-120.
- Freidig, A.P., E.A. Garicano, and F.J.M. Busser. 1998. Estimating Impact of Humic Acid on Bioavailability and Bioaccumulation of Hydrophobic Chemicals in Guppies Using Kinetic Solid-Phase Extraction. *Environmental Toxicology and Chemistry*, 17(6):998-1004.
- Ganf, G.G, and P. Bla• ka. 1974. Oxygen Uptake, Ammonia and Phosphate Excretion by Zooplankton in a Shallow Equatorial Lake (Lake Goerge, Uganda). *Limnol. Oceanog*. 19(2):313-325.

- Gilek, M., M. Björk, D. Broman, N. Kautsky, and C. Näf. 1996. Enhanced Accumulation of PCB Congeners by Baltic Sea Blue Mussels, *Mytilus edulis*, with Increased Algae Enrichment. *Environmental Toxicology and Chemistry*, 15(9):1597-1605.
- Gobas, F.A.P.C. 1993. A Model for Predicting the Bioaccumulation Hydrophobic Organic Chemicals in Aquatic Food-webs: Application to Lake Ontario. *Ecological Modelling*, 69:1-17.
- Gobas, F.A.P.C., E.J. McNeil, L. Lovett-Doust, and G.D. Haffner. 1991. Bioconcentration of Chlorinated Aromatic Hydrocarbons in Aquatic Macrophytes (*Myriophyllum spicatum*). *Environmental Science & Technology*, 25:924-929.
- Gobas, F.A.P.C., Xin Zhang, and Ralph Wells. 1993. Gastrointestinal Magnification: The Mechanism of Biomagnification and Food Chain Accumulation of Organic Chemicals. *Environmental Science & Technology*, 27:2855-2863.
- Gobas, F.A.P.C., M.N. Z-Graggen, X. Zhang. 1995. Time response of the Lake Ontario Ecosystem to Virtual Elimination of PCBs. *Environmental Science & Technology*, 29(8):2038-2046.
- Godshalk, G.L., and J.W. Barko. 1985. Chapter 4, Vegetative Succession and Decomposition in Reservoirs. In D. Gunnison (ed.), *Microbial Processes in Reservoirs*, Dordrecht: Dr. W. Junk Publishers, pp. 59-77.
- Groden, W.T. 1977. *Modeling Temperature and Light Adaptationm of Algae*. Report 2, Cenyer for Ecological Modeling, Rensselaer Polytechnic Institute, Troy, New York, 17 pp.
- Gunnison, D., J.M. Brannon, and R.L. Chen. 1985. Chapter 9, Modeling Geomicrobial Processes in Reservoirs. In D. Gunnison (ed.), *Microbial Processes in Reservoirs*. Dordrecht: Dr. W. Junk Publishers, pp. 155-167.
- Hanna, M. 1990. Evaluation of Models Predicting Mixing Depth. *Can. J. Fish. Aquat. Sci.*, 47:940-947.
- Harris, G.P. 1986. *Phytoplankton Ecology: Structure, Function and Fluctuation*. Chapman and Hall, London, 384 pp.
- Hawker, D.W. and D.W. Connell. 1985. Prediction of Bioconcentration Factors Under Non-Equilibrium Conditions. Chemosphere 14(11/12):1835-1843.
- Hewett, S.W., and B.L. Johnson. 1992. *Fish Bioenergetics 2 Model*. Madison, Wisconsin: University of Wisconsin Sea Grant Institute, 79 pp.
- Hill, I.R., and P.L. McCarty. 1967. Anaerobic Degradation of Selected Chlorinated Pesticides. *Jour. Water Poll. Control Fed.* 39:1259.

- Hill, W.R., and Napolitano, G.E. 1997. PCB Congener Accumulation by Periphyton, Herbivores, and Omnivores. *Archives Environmental Contamination Toxicology*, 32:449-455.
- Hoggan, D.H. 1989. *Computer-Assisted Floodplain Hydrology and Hydraulics*, McGraw-Hill New York, 518 pp.
- Horne, A.J., and C.R. Goldman. 1994. *Limnology* 2nd edition. McGraw-Hill, New York, 576 pp.
- Howick, G.L., F. deNoyelles, S.L. Dewey, L. Mason, and D. Baker. 1993. The Feasibility of Stocking Largemouth Bass in 0.04-ha Mesocosms Used for Pesticide Research. *Environmental Toxicology and Chemistry*, 12:1883-1893.
- Hrbáček, J. 1966. A Morphometrical Study of Some Backwaters and Fish Ponds in Relation to the Representative Plankton Samples. In *Hydrobiological Studies 1*, J. Hrbáček, Ed., Czechoslovak Academy of Sciences, Prague, p. 221-257.
- Hutchinson, G.E. 1957. A Treatise on Limnology, Volume I, Geography, Physics, and Chemistry. John Wiley & Sons, New York, 1015 pp.
- Hutchinson, G.E. 1967. A Treatise on Limnology, Volume II, Introduction to Lake Biology and the Limnoplankton. Wiley & Sons, New York, 1115 pp.
- Imboden, D.M. 1973. Limnologische Transport- und Nährstoffmodelle. *Schweiz. Z. Hydrol.* 35:29-68
- Johanson, R.C., J.C. Imhoff, and H.H. Davis, Jr. 1980. *Users Manual for Hydrological Simulatiuon Program Fortran (HSPF)*. U.S. Environmental Protection Agency, Athens Environmental Research Laboratory, EPA-600/9-80-015, 678 pp.
- Jørgensen, S.E. 1976. A Eutrophication Model for a Lake. Ecol. Modelling, 2:147-165.
- Jørgensen, S.E., H.F. Mejer, M. Friis, L.A. Jørgensen, and J. Hendriksen (Eds.). 1979. *Handbook of Environmental Data and Ecological Parameters*. Copenhagen: International Society of Ecological Modelling.
- Junge, C.O. 1966. Depth distributions for quadratic surfaces and other configurations. In: Hrbáček, J. (Ed.): *Hydrobiological Studies. Vol. 1*, Academia, Prague, pp. 257-265.
- Jupp, B.P., and D.H.N. Spence. 1977a. Limitations on Macrophytes in a Eutrophic Lake, Loch Leven I. Effects of Phytoplankton. *Journal Ecology*, 65:175-186.
- Jupp, B.P., and D.H.N. Spence. 1977b. Limitations on Macrophytes in a Eutrophic Lake, Loch Leven II. Wave Action, Sediments, and Waterfowl Grazing. *Journal Ecology*, 65:431-446.

- Karickhoff, S.W., and K.R. Morris. 1985. Sorption Dynamics of Hydrophobic Pollutants in Sediment Suspensions. *Environmental Toxicology and Chemistry*, 4:469-479.
- Kitchell, J.F., J.F. Koonce, R.V. O'Neill, H.H. Shugart, Jr., J.J Magnuson, and R.S. Booth. 1972. *Implementation of a Predator-Prey Biomass Model for Fishes*. Eastern Deciduous Forest Biome, International Biological Program, Report 72-118. 57 pp.
- Kitchell, J.F., J.F. Koonce, R.V. O'Neill, H.H. Shugart, Jr., J.J Magnuson, and R.S. Booth. 1974. Model of fish biomass dynamics. *Trans. Am. Fish. Soc.* 103:786-798.
- Koelmans, A.A., and E.H.W. Heugens. 1998. Binding Constants of Chlorobenzenes and Polychlorobiphenyls for Algal Exudates. *Water Science Technology*, 37(3):67-73.
- Koelmans, A.A., S.F.M. Anzion, and L. Lijklema. 1995. Dynamics of Organic Micropollutant Biosorption to Cyanobacteria and Detritus. *Environmental Science & Technology*, 29(4):933-940.
- Kremer, J.N., and S.W. Nixon. 1978. *A Coastal Marine Ecosystem*. Springer-Verlag, New York, N.Y., 217 pp.
- Krenkel, P.A., and G.T. Orlob. 1962. Turbulent Diffusion and the Reaeration Coefficient. *Proc. ASCE, Jour. San. Eng. Div.*, 88 (SA 2):53-83.
- Krone, R. B. 1962. Flume Studies of The Transport of Sediment in Estuarial Shoaling Processes: Final Report, Hydraulic Engr. and San. Engr., Research Lab., University of California at Berkeley.
- Lam, R.K., and B.W. Frost. 1976. Model of Copepod Filtering Responses to Changes in Size and Concentration of Food. *Limnol. Oceanogr.* 21:490-500.
- Larsen, D.P., H.T. Mercier, and K.W. Malueg. 1973. Modeling Algal Growth Dynamics in Shagawa Lake, Minnesota, with Comments Concerning Projected Restoration of the Lake. In E.J. Middlebrooks, D.H. Falkenborg, and T.E. Maloney (Eds.). *Modeling the Eutrophication Process*. Logan, Utah: Utah State University, pp. 15-32.
- Le Cren, E.P., and R.H. Lowe-McConnell (Eds.). 1980. *The Functioning of Freshwater Ecosystems*. Cambridge: Cambridge University Press, 588 pp.
- Lehman, J.T., D.B. Botkin, and G.E. Likens. 1975. The Assumptions and Rationales of a Computer Model of Phytoplankton Population Dynamics. *Limnol. and Oceanogr.* 20(3):343-364.

- Leidy, G.R., and R.M. Jenkins. 1977. *The Development of Fishery Compartments and Population Rate Coefficients for Use in Reservoir Ecosystem Modeling*. Contract Rept. CR-Y-77-1, U.S. Army Engineer Waterways Experiment Station, Vicksburg Mississippi, 134 pp.
- Leung, D.K. 1978. *Modeling the Bioaccumulation of Pesticides in Fish*. Report N. 5, Center for Ecological Modeling, Resselaer Polytechnic Institute, Troy, N.Y.
- Liss, P.S., and P.G. Slater. 1974. Flux of Gases Across the Air-Sea Interface. *Nature*, 247:181-184.
- Lyman, W.J., W.F. Reehl, and D.H. Rosenblatt. 1982. *Handbook of Chemical Property Estimation Methods*. McGraw-Hill, New York.
- Maaret, K., K, Leif, and H. Bjarne. 1992. Studies on the Partition Behavior of Three Organic Hydrophobic Pollutants in Natural Humic Water. *Chemosphere*, 24(7):919-925.
- Mabey, W., and T. Mill. 1978. Critical Review of Hydrolysis of Organic Compounds in Water Under Environmental Conditions. *J. Phys. Chem. Ref. Data*, 7:383-415.
- Macek, K.J., M.E. Barrows, R.F. Frasny, and B.H. Sleight III. 1977. Bioconcentration of ¹⁴C-Pesticides by Bluegill Sunfish During Continuous Aqueous Exposure. In *Structure-Activity Correlations in Studies of Toxicity and Bioconcentration with Aquatic Organisms*, G.D. Veith and D. Konasewick, eds.
- Mackay, D., H. Puig, and L.S. McCarty. 1992. An Equation Describing the Time Course and Variability in Uptake and Toxicity of Narcotic Chemicals to Fish. *Environmental Tixcology and Chemistry*, 11:941-951.
- Mancini, J.L. 1983. A Method for Calculating Effects on Aquatic Organisms of Time Varying Concentrations. Water Res. 10:1355-1362.
- Mayer, F. L., Jr., and M. R. Ellersieck. 1986. *Manual of Acute Toxicity: Interpretation and Data Base for 410 Chemicals and 66 Species of Freshwater Animals*: U.S. Department of Interior Fish and Wildlife Service, Resource Publication 160; Wasjington, D.C.
- Mayio, A.E., and G.H. Grubbs. 1993. Nationwide Water-Quality Reporting to the Congress as Required Under Section 305(b) of the Clean Water Act. In *National Water Summary 1990-91*, Water Supply Paper 2400; Washington, D.C.: U.S. Geological Survey, pp. 141-146.
- McCarty, L.S., G.W. Ozburn, A.D. Smith, and D.G. Dixon. 1992. Toxicokinetic Modeling of Mixtures of Organic Chemicals. *Environmental Toxicology and Chemistry*, 11:1037-1047.
- McIntire, C.D. 1968. Structural Characteristics of Benthic Algal Communities in Laboratory Streams. *Ecology* 49(3):520-537.

- McIntire, C.D. 1973. Periphyton Dynamics in Laboratory Streams: a Simulation Model and Its Implications. *Ecological Monographs* 43(3):399-419.
- McIntire, C.D., and J.A. Colby. 1978. A Hierarchical Model of Lotic Ecosystems. *Ecological Monographs* 48:167-190.
- McKay, M.D., W.J. Conover, and R.J. Beckman. 1979. A Comparison of Three Methods for Selecting Values of Input Variables in the Analysis of Output from a Computer Code. *Technometrics* 211:239-245.
- McKim, J.M., G.F. Olson, G.W. Holcombe, and E.P. Hunt. 1976. Long-Term Effects of Methylmercuric Chloride on Three Generations of Brook Trout (*Salvelinus fontinalis*): Toxicity, Accumulation, Distribution, and Elimination. *Journal Fisheries Research Board Canada*, 33(12):27226-2739.
- McKim, J.M., P. Schmeider, and G. Veith. 1985. Absorption Dynamics of Organic Chemical Transport Across Trout Gills as Related to Octanol-Water Partition Coefficient. *Toxicology and Applied Pharmacology*, 77:1-10.
- Megard, R.O., W.S. Comles, P.D. Smith, and A.S. Knoll. 1979. Attenuation of Light and Daily Integral Rates of Photosynthesis Attained by Planktonic Algae. *Limnol. Oceanogr.*, 24:1038-1050.
- Muirhead-Thomson, R.C. 1987. Pesticide Impact on Stream Fauna with Special Reference to Macroinvertebrates. Cambridge: Cambridge University Press, 275 pp.
- Mullin, M.M. 1963. Some Factors Affecting the Feeding of Marine Copepods of the Genus *Calanus*. *Limnol. Oceanogr.* 8:239-250.
- Mullin, M.M., E.F. Stewart, and F.J. Foglister. 1975. Ingestion by Planktonic Grazers as a Function of Concentration of Food. *Limnol. Oceanog.* 20:259-262.
- Nalewajko, C. 1966. Photosynthesis and Excretion in Various Planktonic Algae. *Limnol. Oceanogr.*, 11:1-10.
- Nichols, D.S., and D.R. Keeney. 1976. Nitrogen Nutrition of *Myriphyllum spicatum*: Uptake and Translocation of ¹⁵N by Shoots and Roots. *Freshwater Biology* 6:145-154.
- O'Connor, D.J., and J.P. Connolly. 1980. The Effect of Concentration of Adsorbing Solids on the Partition Coefficient. *Water Research*, 14:1517-1523.
- O'Connor, D.J., and W.E. Dobbins. 1958. Mechanism of Reaeration in Natural Streams. *ASCE Transactions*, pp. 641-684, Paper No. 2934.

- O'Connor, D.J., J.L. Mancini, and J.R. Guerriero. 1981. Evaluation of Factors Influencing the Temporal Variation of Dissolved Oxygen in the New York Bight, Phase II. Manhattan College, Bronx, New York
- Odum, E.P., and A.A. de la Cruz. 1963. Detritus as a Major Component of Ecosystems. *Amer. Inst. Biol. Sci. Bull.*, 13:39-40.
- Oliver, B.G., and A.J. Niimi. Trophodynamic Analysis of Polychlorinated Biphenyl Congeners and Other Chlorinated Hydrocarbons in the Lake Ontario Ecosystem. *Environ. Sci. Technol.*, 22(4):388-397.
- O'Neill, R.V. 1969. Indirect Estimation of Energy Fluxes in Animal Food Webs. *Jour. Theoret. Biol.*, 22:284-290.
- O'Neill, R.V., D.L. DeAngelis, J.B. Waide, and T.F.H. Allen. 1986. *A Hierarchical Concept of the Ecosystem*. Princeton University Press, Princeton, N.J.
- O'Neill, R.V., R.A. Goldstein, H.H. Shugart, and J.B. Mankin. 1972. *Terrestrial Ecosystem Energy Model*. Eastern Deciduous Forest Biome, International Biological Program Report 72-19.
- Owens, M., R.W. Edwards, and J.W. Gibbs. 1964. Some Reaeration Studies ion Streams. *Internat. Jour. Air Water Poll.* 8:469-486.
- Palisade Corporation. 1991. *Risk Analysis and Simulation Add-In for Lotus 1-2-3*. Newfield New York, 342 pp.
- Park, R.A. 1978. *A Model for Simulating Lake Ecosystems*. Center for Ecological Modeling Report No. 3, Rensselaer Polytechnic Institute, Troy, New York, 19 pp.
- Park, R.A. 1984. TOXTRACE: A Model to Simulate the Fate and Transport of Toxic Chemicals in Terrestrial and Aquatic Environments. *Acqua e Aria*, No. 6, p. 599-607 (in Italian).
- Park, R.A. 1990. *AQUATOX*, a Modular Toxic Effects Model for Aquatic Ecosystems. Final Report, EPA-026-87; U.S. Environmental Protection Agency, Corvallis, Oregon.
- Park, R.A. 1999. Evaluation of AQUATOX for Predicting Bioaccumulation of PCBs in the Lake Ontario Food Web. In: AQUATOX for Windows: A Modular Fate and Effects Model for Aquatic Ecosystems–Volume 3: Model Validation Reports. U.S. Environmental Protection Agency 2000. EPA-823-R-00-008
- Park, R.A., J.J. Anderson, G.L. Swartzman, R. Morison, and J.M. Emlen. 1988. Assessment of Risks of Toxic Pollutants to Aquatic Organisms and Ecosystems Using a Sequential Modeling Approach. In *Fate and Effects of Pollutants on Aquatic Organisms and Ecosystems*, 153-165. EPA/600/9-88/001. Athens, Ga.: U.S. Environmental Protection Agency

- Park, R.A., and C.D. Collins. 1982. Realism in Ecosystem Models. *Perspectives in Computing* 2(2):18–27.
- Park, R.A., C.D. Collins, C.I. Connolly, J.R. Albanese, and B.B. MacLeod. 1980. *Documentation of the Aquatic Ecosystem Model MS.CLEANER, A Final Report for Grant No. R80504701*, U.S. Environmental Protection Agency, Environmental Research Laboratory, Athens, Georgia. 112 pp.
- Park, R.A., C.D. Collins, D.K. Leung, C.W. Boylen, J.R. Albanese, P. deCaprariis, and H. Forstner. 1979. The Aquatic Ecosystem Model MS.CLEANER. In *State-of- the-Art in Ecological Modeling*, edited by S.E. Jorgensen, 579–602. International Society for Ecological Modelling, Denmark.
- Park, R.A., C.I. Connolly, J.R. Albanese, L.S. Clesceri, G.W. Heitzman, H.H. Herbrandson, B.H. Indyke, J.R. Loehe, S. Ross, D.D. Sharma, and W.W. Shuster. 1980. *Modeling Transport and Behavior of Pesticides and Other Toxic Organic Materials in Aquatic Environments*. Center for Ecological Modeling Report No. 7. Rensselaer Polytechnic Institute, Troy, New York. 163 pp.
- Park, R.A., C.I. Connolly, J.R. Albanese, L.S. Clesceri, G.W. Heitzman, H.H. Herbrandson, B.H. Indyke, J.R. Loehe, S. Ross, D.D. Sharma, and W.W. Shuster. 1982. *Modeling the Fate of Toxic Organic Materials in Aquatic Environments*. U.S. Environmental Protection Agency Rept. EPA–600/S3–82–028, Athens, Georgia.
- Park, R.A., T.W. Groden, and C.J. Desormeau. 1979. Modifications to the Model CLEANER Requiring Further Research. In *Perspectives on Lake Ecosystem Modeling*, edited by D. Scavia and A. Robertson. Ann Arbor Science Publishers, Inc., 22 pp.
- Park, R.A., B.H. Indyke, and G.W. Heitzman. 1981. Predicting the Fate of Coal-Derived Pollutants in Aquatic Environments. Paper presented at Energy and Ecological Modelling symposium, Louisville, Kentucky, April 2023, 1981. Developments in Environmental Modeling 1. 7 pp.
- Park, R.A., B.B. MacLeod, C.D. Collins, J.R. Albanese, and D. Merchant. 1985. *Documentation of the Aquatic Ecosystem MINI.Cleaner, A Final Report for Grant No. R806299020*. U.S. Environmental Protection Agency, Environmental Research Laboratory, Athens, Georgia. 85 pp.
- Park, R.A., R.V. O'Neill, J.A. Bloomfield, H.H. Shugart, Jr., R.S. Booth, J.F. Koonce, M.S. Adams, L.S. Clesceri, E.M. Colon, E.H. Dettman, R.A. Goldstein, J.A. Hoopes, D.D. Huff, S. Katz, J.F. Kitchell, R.C. Kohberger, E.J. LaRow, D.C, McNaught, J.L. Peterson, D. Scavia, J.E. Titus, P.R. Weiler, J.W. Wilkinson, and C.S. Zahorcak. 1974. A Generalized Model for Simulating Lake Ecosystems. Simulation, 23(2):30–50. Reprinted in Benchmark Papers in Ecology.

- Park, R.A., D. Scavia, and N.L. Clesceri. 1975. CLEANER, The Lake George Model. In *Ecological Modeling in a Management Context*. Resources for the Future, Inc., Washington, D.C.
- Parker, R.A. 1972. Estimation of Aquatic Ecosystem Parameters. *Verh. Internat. Verein. Limnol.* 18:257-263.
- Parsons, T.R., R.J. LeBresseur, J.D. Fulton, and O.D. Kennedy. 1969. Production Studies in the Strait of Georgia II. Secondary Production Under the Fraser River Plume, February to May, 1967. *Jour. Exp. Mar. Biol. Ecol.* 3:39-50.
- Partheniades, E. 1965. Erosion and Deposition of Cohesive Soils. *ASCE Jour. Hydrol. Div.* pp. 105-138.
- Partheniades, E. 1971. "Erosion and Deposition of Cohesive Materials". In River Mechanics, H. W. Shen Ed. Chapter 25. Water Resources Publications, Littleton, Colorado.
- Patten, B.C., D.A. Egloff, and T.H. Richardson. 1975. Total Ecosystem Model for a Cove in Lake Texoma. In B.C. Patten (Ed.) *Systems Analysis and Simulation in Ecology. Vol. III.* New York: Academic Press, pp. 205-241.
- Press, W.H., B.P. Flannery, S.A. Teukolsky, and W.T. Vetterling. 1986. Numerical Recipes: The Art of Scientific Computing. Cambridge University Press, Cambridge, U.K. 818 pp.
- Redfield, A.C. 1958. The Biological Control of Chemical Factors in the Environment. *American Scientist* 46:205-222.
- Riley, G.A. 1963. Theory of Food-Chain Relations in the Ocean. *The Sea*, 2.
- Sand-Jensen, K. 1977. Effects of Epiphytes on Eelgrass (*Zostera marina* L.) in Danish Coastal Waters. *Marine Technology Society Journal* 17:15-21.
- Saunders, G.W. 1980. 7. Organic Matter and Decomposers. In E.P. Le Cren and R.H. Lowe-McConnell (Eds.), *The Functioning of Freshwater Ecosystems*. Cambridge: Cambridge University Press, pp. 341-392.
- Scavia, D. 1979. Chapter 6 The Use of Ecological Models of Lakes in Synthesizing Available Information and Identifying Research Needs. In D. Scavia and A. Robertson (Eds.) *Perspectives on Lake Ecosystem Modeling*. Ann Arbor, Michigan: Ann Arbor Science, pp. 109-168.
- Scavia, D. 1980. An Ecological Model of Lake Ontario. *Ecological Modelling* 8:49-78.

- Scavia, D., B.J. Eadie, and A. Robertson. 1976. *An Ecological Model for Lake Ontario—Model Formulation, Calibration, and Preliminary Evaluation*. Tech. Report ERL 371-GLERL 12, National Oceanic and Atmospheric Administration, Boulder, Colorado.
- Scavia, D., and R.A. Park. 1976. Documentation of Selected Constructs and Parameter Values in the Aquatic Model CLEANER. *Ecological Modelling* 2(1):33–58.
- Schwarzenbach, R.P., P.M. Gschwend, and D.M. Imboden. 1993. *Environmental Organic Chemistry*. Wiley and Sons, Inc., New York.
- Sedell, J.R., F.J. Triska, and N.S. Triska. 1975. The Processing of Conifer and Hardwood Leaves in Two Coniferous Forest Streams: I. Weight Loss and Associated Invertebrates. *Herh. Internat. Verein. Limnol.*, 19:1617-1627.
- Sijm, D.T.H.M., K.W. Broersen, D.F de Roode, and P. Mayer. 1998. Bioconcentration Kinetics of Hydrophobic Chemicals in Different Densities of *Chlorella Opyrenoidosa*. *Environmental Toxicology and Chemistry* 17:9:1695-1704.
- Skoglund, R.S., K. Stange, and D.L. Swackhamer. 1996. A Kinetics Model for Predicting the Accumulation of PCBs in Phytoplankton. *Environmental Science and Technology* 30:7:2113-2120.
- Smayda, T.J. 1971. Some Measurements of the Sinking Rate of Fecal Pellets. *Limnology and Oceanography* 14:621-625.
- Smayda, T.J. 1974. Some Experiments on the Sinking Characteristics of Two Freshwater Diatoms. *Limnology and Oceanography* 19:628-635.
- Smejtek, P., and S. Wang. 1993. Distribution of Hydrophobic Ionizable Xenobiotics Between Water and Lipid Membranes: Pentachlorophenol and Pentachlorophenate. A Comparison with Octanol-Water Partition. *Archives of Environmental Contamination and Toxicology*, 25(3):394.
- Smith, D.J. 1978. WQRRS, Generalized Computer Program for River-Reservoir Systems. U.S. Army Corps of Engineers, Hydrologic Engineering Center (HEC), Davis, California Users Manual 401-100, 100A, 210 pp.
- Southworth, G.R., J.J. Beauchamp, and P.K. Schmieder. 1978. Bioaccumulation Potential of Polycyclic Aromatic Hydrocarbons in *Daphnia pulex*. *Water Res.*, 12:973-977.
- Spacie, A., and J.L. Hamelink. 1982. Alternative Models for Describing the Bioconcentration of Organics in Fish. *Environmental Toxicology and Chemistry*, 1:309-320.

- Stange, K., and D.L. Swackhamer. 1994. Factors Affecting Phytoplankton Species-Specific Differences in Accumulation of 40 Polychlorinated Biphenyls (PCBs). *Environmental Toxicology and Chemistry*, 13(11):1849-1860.
- Steele, J.H. 1962. Environmental Control of Photosynthesis in the Sea. *Limnol. Oceanogr.*, 7:137-150.
- Steele, J.H. 1974. *The Structure of Marine Ecosystems*. Harvard University Press, Cambridge, Massachusetts, 128 pp.
- Steele, J.H., and M.M. Mullin. 1977. Zooplankton Dynamics. In E.D. Goldberg, I.N. McCave, J.J. O'Brien, and J.H. Steele (Eds.), *The Sea Vol. 6: Marine Modeling*, New York: Wiley-Interscience, p. 857.
- Stefan, H.G., and X. Fang. 1994. Dissolved Oxygen Model for Regional Lake Analysis. *Ecological Modelling* 71:37-68.
- Stewart, D.C. 1975. *Mathematical Modelling of the Ecosystem of Lough Neagh*. Ph.D. Dissertation, Queen's University, Belfast, Northern Ireland.
- Straškraba, M. 1973. Limnological Basis for Modeling Reservoir Ecosystems. In Ackermann, W.C., G.F. White, and E.B. Worhtington (eds.) *Man-Made Lakes: Their Problems and Environmental Effects*. Geophys. Momogr. Series Vol. 17, London, pp. 517-538.
- Straškraba, M. and A.H. Gnauck. 1985. *Freshwater Ecosystems: Modelling and Simulation*. Developments in Environmental Modelling, 8. Elsevier Science Publishers, Amsterdam, The Netherlands. 309 pp.
- Stumm, W., and J.J. Morgan. 1996. *Aquatic Chemistry: Chemical Equilibria and Rates in Natural Waters 3rd Edition*. New York: John Wiley & Sons, 1022 pp.
- Suárez, L.A., and M.C. Barber. 1992. PIRANHA Version 2.0, FGETS Version 3.0-11 User's Manual, In *PIRANHA Pesticide and Industrial Chemical Risk Analysis and Hazard Assessment*. Athens, Georgia: U.S. Environmental Protection Agency.
- Suter, G.W., II, A.E. Rosen, and E. Linder. 1986. 4. Analysis of Extrapolation Error. *User's Manual for Ecological Risk Assessment*. Oak Ridge National Laboratory, ORNL-6251, pp. 49-81.
- Swackhamer, D.L., and R.S. Skoglund. 1991. The Role of Phytoplankton in the Partitioning of Hydrophobic Organic Contaminants in Water. In Baker, R.A., ed., *Organic Substances and Sediments in Water Vol. 2 C Processes and Analytical*, Lewis: Chelsea MI, pp. 91-105.
- Swackhamer, D.L., and R.S. Skoglund. 1993. Bioaccumulation of PCBs by Algae: Kinetics versus Equilibrium. *Environmental Toxicology & Chemistry*, 12:831-838.

- Thomann, R.V. 1989. Bioaccumulation Model of Organic Chemical Distribution in Aquatic Food Chains. *Environmental Science & Technology*, 23:699-707.
- Thomann, R.V., and J.J. Fitzpatrick. 1982. *Calibration and Verification of a Mathematical Model of the Eutrophication of the Potomac Estuary*. Prepared for Department of Environmental Services, Government of the District of Columbia, Washington, D.C.
- Thomann, R.V., D.M. Di Toro, R.P. Winfield, and D.J. O'Connor. 1975. *Mathematical Modeling of Phytoplankton in Lake Ontario, Part 1. Model Development and Verification*. Manhattan College, Bronx, New York, for U.S. Environmental Protection Agency EPA-600/3-75-005.
- Thomann, R.V., and J.A. Mueller. 1987. *Principles of Surface Water Quality Modeling and Control*, Harper Collins: new York N.Y., 644 pp.
- Thomann, R.V., J.A. Mueller, R.P. Winfield, and C.-R. Huang. 1991. Model of Fate and Accumulation of PCB Homologues in Hudson Estuary. *Jour. Environ. Engineering*, 117(2):161-178.
- Thomann, R.V., J. Segna, and R. Winfield. 1979. *Verification Analysis of Lake Ontario and Rochester Embayment Three-Dimensional Eutrophication Models*. Manhattan College, Bronx, New York, for U.S. Environmental Protection Agency.
- Titus, J.E., M.S. Adams, P.R. Weiler, R.V. O'Neill, H.H. Shugart, Jr., and J.B. Mankin. 1972. *Production Model for* Myriophyllum spicatum *L.* Memo Rept. 72-19, U.S. International Biological Program Eastern Deciduous Forest Biome, University of Wisconsin, Madison, 17 pp.
- Toetz, D.W. 1967. The Importance of Gamete Losses in Measurements of Freshwater Fish Production. *Ecology*. 48:1017-1020.
- US EPA. 1988. The Effects of Chloropyrifos on a Natural Aquatic System: A Research Design for Littoral Enclosure Studies and Final Research Report. U.S. Environmental Protection Agency, Environmental Research Laboratory, Duluth, Minnessota, 194 pp.
- US EPA. 1991. *Hydrological Simulation Program FORTRAN User's Manual for Release 10* (Pre-release Draft Version). U.S. EPA Technology Development and Applications Branch in cooperation with USGS Water Resources Division, Office of Surface Water. By Bicknell, B.R., J.C. Imhoff, J.L. Kittle, A.S. Donigian, and -R.C. Johanson.
- US EPA. 1995. Great Lakes Water Quality Initiative Technical Support Document for the Procedure to Determine Bioaccumulation Factors. EPA-820-B-95-005, U.S. Environmental Protection Agency, Washington, D.C.

- US EPA. 1998. *National Water Quality Inventory: 1996 Report to Congress*. EPA841-R-97-008. April 1998.
- Verduin, 1982. Components Contributing to Light Extinction in Natural Waters: Method of Isolation. *Arch. Hydrobiol.*, 93(3):303-312.
- Ward 1963, ASCE 1989, 6:1-16
- Watt, W.D. 1966. Release of Dissolved Organic Material From the Cells of Phytoplankton Species in Natural and Mixed Populations. *Proceedings of the Royal Society, London*, B 164:521-525.
- Weininger, D. 1978. *Accumulation of PCBs by Lake Trout in Lake Michigan*. Ph.D. Dissertation, University of Wisconsin, Madison, 232 pp.
- Wetzel, R.G. 1975. Limnology, W.B. Saunders, Philadelphia, 743 pp.
- Wetzel, R.G., P.H. Rich, M.C. Miller, and H.L. Allen. 1972. Metabolism of Dissolved and Particulate Detrital Carbon in a Temperate Hard-water Lake. *in* U. Melchiorri-Santolinii and J.W. Hopton (eds.) *Detritus and Its Role in Aquatic Ecosystems, Mem. Ist. Ital. Idobiol.*, 29(Suppl):185-243.
- Westlake, D.F. 1967. Some Effects of Low Velocity Currents on the Metabolism of Aquatic Macrophytes. *Journal Experimental Botany* 18:187-205.
- Whitman, W.G. 1923. The two-film theory of gas absorption. *Chem. Metal. Eng.* 29:146-148.
- Wood, L.W., P. O.Keefe, and B. Bush. 1997. Similarity Analysis of PAH and PCB Bioaccumulation Patterns in Sediment-Exposed *Chironomus tentans* Larvae. *Environmental Toxicology and Chemistry*, 16(2):283-292.

APPENDIX A. GLOSSARY OF TERMS

Taken in large part from: The Institute of Ecology. 1974. *An Ecological Glossary for Engineers and Resource Managers*. TIE Publication #3, 50 pp.

Abiotic nonliving, pertaining to physico-chemical factors only

Adsorption the adherence of substances to the surfaces of bodies with which they are in

contact

Aerobic living, acting, or occurring in the presence of oxygen

Algae any of a group of chlorophyll-bearing aquatic plants with no true leaves,

stems, or roots

Allochthonous material derived from outside a habitat or environment under consideration

Algal bloom rapid and flourishing growth of algae

Alluvial of alluvium

Alluvium sediments deposited by running water

Ambient surrounding on all sides

Anaerobic capable of living or acting in the absence of oxygen
Anoxic pertaining to conditions of oxygen deficiency
Aphotic below the level of light penetration in water

Assimilation transformation of absorbed nutrients into living matter

Autochthonous material derived from within a habitat, such as through plant growth

Benthic pertaining to the bottom of a water body; pertaining to organisms that live on

the bottom

Benthos those organisms that live on the bottom of a body of water

Biodegradable can be broken down into simple inorganic substances by the action of

decomposers (bacteria and fungi)

Biochemical oxygen the amount of oxygen required to decompose a given amount of organic

demand (BOD) matter

Biomagnification the step by step concentration of chemicals in successive levels of a food

chain or food web

Biomass the total weight of matter incorporated into (living and/or dead) organisms

Biota the fauna and flora of a habitat or region **Chlorophyll** the green, photosynthetic pigments of plants

Colloid a dispersion of particles larger than small molecules and that do not settle out

of suspension

Consumer an organism that consumes another

Copepods a large subclass of usually minute, mostly free-swimming aquatic crustaceans

Crustacean a large class of arthropods that bear a horny shell bacteria and fungi that break down organic detritus

Detritus dead organic matter

Diatom any of class of minute algae with cases of silica

Diurnal pertaining to daily occurrence

Dynamic a state of relative balance between processes having opposite effects

equilibrium

Ecology the study of the interrelationships of organisms with and within their

environment

Ecosystem a biotic community and its (living and nonliving) environment considered

together

Emergent aquatic plants, usually rooted, which have portions above water for part of

their life cycle

Environment the sum total of all the external conditions that act on an organism the well mixed surficial layer of a lake; above the hypolimnion

Epiphytes plants that grow on other plants, but are not parasitic

Equilibrium a steady state in a dynamic system, with outflow balancing inflow

Euphotic pertaining to the upper layers of water in which sufficient light penetrates to

permit growth of plants

Eutrophic aquatic systems with high nutrient input and high plant growth

Fauna the animals of a habitat or region

Flood plain that part of a river valley that is covered in periods of high (flood) water

Flora plants of a habitat or region Fluvial pertaining to a stream

Food chain animals linked by linear predator-prey relationships with plants or detritus at

the base

Food web similar to food chain, but implies cross connections

Forage fish fish eaten by other fish

Habitat the environment in which a population of plants or animals occurs **Humic** pertaining to the partial decomposition of leaves and other plant material

Hydrodynamics the study of the movement of water

Hypolimnion the lower layer of a stratified water body, below the well mixed zone

Influent anything flowing into a water body

Inorganic pertaining to matter that is neither living nor immediately derived from living

matter

Invertebrate animals lacking a backbone

Limiting factor an environmental factor that limits the growth of an organism; the factor that

is closest to the physiological limits of tolerance of that organism

Limnetic zone the open water zone of a lake or pond from the surface to the depth of

effective light penetration

Limnology the study of inland waters

Littoral zone the shoreward zone of a water body in which the light penetrates to the

bottom, thus usually supporting rooted aquatic plants

Macrofauna animals visible to the naked eye

Macrophytes large (non-microscopic), usually rooted, aquatic plants

Nutrients chemical elements essential to life

Omnivorous feeding on a variety of organisms and organic detritus

Organic chemical compounds containing carbon;

Overturn the complete circulation or mixing of the upper and lower waters of a lake

when temperatures (and densities) are similar

Oxygen depletion exhaustion of oxygen by chemical or biological use

Parameter a measurable, variable quantity as distinct from a statistic

Pelagic zone open water with no association with the bottom

Periphyton community of algae and associated organisms, usually small but densely set,

closely attached to surfaces on or projecting above the bottom

Oxidation a reaction between molecules, ordinarily involves gain of oxygen

Photic zone the region of aquatic environments in which the intensity of light is sufficient

for photosynthesis

Phytoplankton small, mostly microscopic algae floating in the water column

Plankton small organisms floating in the water

Pond a small, shallow lake

Population a group of organisms of the same species

Predator an organism, usually an animal, that kills and consumes other organisms

Prey an organism killed and at least partially consumed by a predator

Producer an organism that can synthesize organic matter using inorganic materials and

an external energy source (light or chemical)

Production the amount of organic material produced by biological activity

Productivity the rate of production of organic matter

Productivity, the rate of production by plants

primary

Productivity, the rate of production by consumers

secondary

Reservoir an artificially impounded body of water

Riverine pertaining to rivers

Rough fish a non-sport fish, usually omnivorous in food habits

Sediment any mineral and/or organic matter deposited by water or air

Siltation the deposition of silt-sized and clay-sized (smaller than sand-sized) particles **Stratification** division of a water body into two or more depth zones due to temperature or

density

Substrate the layer on which organisms grow; the organic substance attacked by

decomposers

Successionthe replacement of one plant assemblage with another through timeTolerancean organism's capacity to endure or adapt to unfavorable conditionsTrophic levelall organisms that secure their food at a common step in the food chainTurbiditycondition of water resulting from suspended matter, including inorganic and

organic material and plankton

Volatilization the act of passing into a gaseous state at ordinary temperatures and pressures

Wastewater water derived from a municipal or industrial waste treatment plant

Wetlands land saturated or nearly saturated with water for most of the year; usually

vegetated

Zooplankton

small aquatic animals, floating, usually with limited swimming capability

APPENDIX B. USER-SUPPLIED PARAMETERS AND DATA

The model has many parameters and internal variables. Most of these are linked to data structures such as ChemicalRecord, SiteRecord, and ReminRecord, which in turn may be linked to input forms that the user accesses through the Windows environment. Although consistency has been a goal, some names may differ between the code and documentation

INTERNAL	TECH DOC	DESCRIPTION	UNITS
ChemicalRecord	Chemical Underlying Data	For each Chemical Simulated, the following parameters are required	
	Data	parameters are required	
ChemName	N/A	Chemical's Name. Used for Reference only.	N/A
CASRegNo	N/A	CAS Registry Number. Used for Reference only.	N / A
MolWt	MolWt	molecular weight of pollutant	(g/mol)
Solubility	N / A	Not utilized as a parameter by the code.	(ppm)
Henry	Henry	Henry's law constant	(atm m3 mol-1)
pka	pKa	acid dissociation constant	negative log
VPress	N/A	Not utilized as a parameter by the code.	mm Hg
LogP	LogKow	log octanol-water partition coefficient	(unitless)
En	En	Arrhenius activation energy	(cal/mol)
KMDegrdn	MicrobialDegrdn	rate of loss due to microbial degradation	(µg/L d)
KMDegrAnaerobic	KAnaerobic	decomposition rate at 0 g/m3 oxygen	(1/d)
KUnCat	KUncat	the measured first-order reaction rate at pH 7	(1/d)
KAcid	KAcidExp	pseudo-first-order acid-catalyzed rate constant for a given pH	(1/d)
KBase	KBaseExp	pseudo-first-order rate constant for a given pH	(1/d)
PhotolysisRate	KPhot	direct photolysis first-order rate constant	(1/d)
OxRateConst	N/A	Not utilized as a parameter by the code.	(L/ mol d)
KPSed	KPSed	detritus-water partition coefficient	(L/kg)
Weibull_Shape	Shape	parameter expressing variability in toxic response	(unitless)
ChemIsBase		if the compound is a base	(True/False)

INTERNAL	TECH DOC	DESCRIPTION	UNITS
SiteRecord	Site Underlying Data	For each Segment Simulated, the following parameters are required	
SiteName	N/A	Site's Name. Used for Reference only.	N/A
ECoeffWater	ExtinctH2O	light extinction of wavelength 312.5 nm in pure water	(1/m)
SiteLength	Length	maximum effective length for wave setup	(m)
Volume	Volume	initial volume of site	(m3)
Area	Area	site area	(m2)
ZMean	ZMean	mean depth	(m)
ZMax	ZMax	maximum depth	(m)
TempMean	TempMean	mean annual temperature	(°C)
TempRange	TempRange	annual temperature range	(°C)
Latitude	Latitude	latitude	(°, decimal)
LightMean	LightMean	mean annual light intensity	(ly/d)
LightRange	LightRange	annual range in light intensity	(ly/d)
AlkCaCO3	N / A	Not utilized as a parameter by the code.	mg/L
HardCaCO3	N / A	Not utilized as a parameter by the code.	mg CaCO3 / L
SO4Conc	N / A	Not utilized as a parameter by the code.	mg/L
TotalDissSolids	N / A	Not utilized as a parameter by the code.	mg/L
StreamType	Stream Type	concrete channel, dredged channel, natural channel	Choice from List
Channel_Slope	Slope	slope of channel	(m/m)
Max_Chan_Depth	Max_Chan_Depth	depth at which flooding occurs	(m)
SedDepth	SedDepth	maximum sediment depth	(m)
LimnoWallArea	LimnoWallArea	area of limnocorral walls; only relevant to limnocorral	(m2)
MeanEvap	MeanEvap	mean annual evaporation	inches / year
UseEnteredManning		do not determine Manning coefficient from streamtype	(true/false)
EnteredManning	Manning	manually entered Manning coefficient.	s / m1/3
ReminRecord	Remineralization Data	For each simulation, the following parameters are required	

INTERNAL	TECH DOC	DESCRIPTION	UNITS
DecayMax_Lab	DecayMax	maximum decomposition rate	(g/g·d)
Q10	NA	Not utilized as a parameter by the code.	(unitless)
TOpt	TOpt	optimum temperature	(°C)
TMax	TMax	maximum temperature tolerated	(°C)
TRef	NA	Not utilized as a parameter by the code.	(°C)
pHMin	pHMin	minimum pH below which limitation on biodegradation rate occurs.	рН
рНМах	pHMax	maximum pH above which limitation on biodegradation rate occurs.	pН
Org2Phosphate	Org2Phosphate	ratio of phosphate to organic matter (unitless)	(unitless)
Org2Ammonia	Org2Ammonia	ratio of ammonia to organic matter	(unitless)
O2Biomass	O2Biomass	ratio of oxygen to organic matter	(unitless)
O2N	O2N	ratio of oxygen to nitrogen	(unitless)
KSed	KSed	intrinsic settling rate	(m/d)
PSedRelease	N / A	Not utilized as a parameter by the code.	(g/m2·d)
NSedRelease	N / A	Not utilized as a parameter by the code.	(g/m2·d)
DecayMax_Refr	ColonizeMax	maximum colonization rate under ideal conditions	(g/g·d)
ZooRecord	Animal Underlying Data	For each animal in the simulation, the following parameters are required	
AnimalName	N/A	Animal's Name. Used for Reference only.	N/A
FHalfSat	FHalfSat	half-saturation constant for feeding by a predator	(g/m3)
CMax	CMax	maximum feeding rate for predator	(g/g·d)
BMin	BMin	minimum prey biomass needed to begin feeding	(g/m3)
Q10	Q10	slope or rate of change per 10°C temperature change	(unitless)
TOpt	TOpt	optimum temperature	(°C)
TMax	TMax	maximum temperature tolerated	(°C)
TRef	TRef	adaptation temperature below which there is no acclimation	(°C)
EndogResp	EndogResp	basal respiration rate at 0° C for given predator	(1/day)

INTERNAL	TECH DOC	DESCRIPTION	UNITS
KResp	KResp	proportion assimilated energy lost to specific dynamic action	(unitless)
KExcr	KExcr	proportionality constant for excretion:respiration	(unitless)
PctGamete	PctGamete	fraction of adult predator biomass that is in gametes	(unitless)
GMort	GMort	gamete mortality	(1/d)
KMort	KMort	intrinsic mortality rate	(g/g·d)
KCap	KCap	carrying capacity	(mg/L)
MeanWeight	WetWt	mean wet weight of organism	(g)
FishFracLipid	LipidFrac	fraction of lipid in organism	(g lipid/g organ
LifeSpan	LifeSpan	mean lifespan in days	days
Animal_Type	Animal Type	Animal Type (Fish, Pelagic Invert, Benthic Invert, Benthic Insect)	Choice from List
AveDrift	Dislodge	fraction of biomass subject to drift per day	fraction / day
AutoSpawn		Does AQUATOX calculate Spawn Dates	(true/false)
SpawnDate13		Automatically Entered Spawn Dates	(date)
UnlimitedSpawning		Allow fish to spawn unlimited times each year	(true/false)
SpawnLimit		Number of spawns allowed for this species this year	(integer)
UseAllom_C		Use Allometric Consumption Equation	(true/false)
CA		Allometric Consumption Parameter	(real number)
СВ		Allometric Consumption Parameter	(real number)
UseAllom_R		Use Allometric Consumption Respiration	(true/false)
RA		Allometric Respiration Parameter	(real number)
RB		Allometric Respiration Parameter	(real number)
UseSet1		Use "Set 1" of Allometric Respiration Parameters	(true/false)
RQ		Allometric Respiration Parameter	(real number)
RK1		Allometric Respiration Parameter	(real number)

INTERNAL	TECH DOC	DESCRIPTION	UNITS
PlantRecord	Plant Underlying Data	For each Plant in the Simulation, the following parameters are required	
PlantName		Plant's Name. Used for Reference only.	N/A
PlantType	Plant Type	Plant Type: (Phytoplankton, Periphyton, Macrophytes)	Choice from List
LightSat	LightSat	light saturation level for photosynthesis	(ly/d)
KPO4	KP	half-saturation constant for phosphorus	(gP/m3)
KN	KN	half-saturation constant for nitrogen	(gN/m3)
KCarbon	KCO2	half-saturation constant for carbon	(gC/m3)
Q10	Q10	slope or rate of change per 10°C temperature change	(unitless)
TOpt	TOpt	optimum temperature	(°C)
TMax	TMax	maximum temperature tolerated	(°C)
TRef	TRef	adaptation temperature below which there is no acclimation	(°C)
PMax	PMax	maximum photosynthetic rate	(1/d)
KResp	KResp	coefficient of proportionality btwn. excretion and photosynthesis at optimal l	(unitless)
KMort	KMort	intrinsic mortality rate	(g/g·d)
EMort	EMort	exponential factor for suboptimal conditions	(unitless)
KSed	KSed	intrinsic settling rate	(m/d)
ESed	ESed	exponential settling coefficient	(unitless)
UptakePO4	Uptake Phosphorus	fraction of photosynthate that is nutrient	(unitless)
UptakeN	Uptake Nitrogen	fraction of photosynthate that is nutrient	(unitless)
ECoeffPhyto	EcoeffPhyto	attenuation coefficient for given alga	(1/m-g/m3)
CarryCapac	KCap	carrying capacity of periphyton	(g/m2)
Red_Still_Water	RedStillWater	reduction in photosynthesis in absence of current	(unitless)
Macrophyte_Type	Macrophyte Type	Type of macrophyte (benthic, rooted floating, free-floating)	Choice from List

INTERNAL	TECH DOC	DESCRIPTION	UNITS
AnimalToxRecord	Animal Toxicity Parameters	For each Chemical Simulated, the following parameters are required for each an	
LC50	LC50	external concentration of toxicant at which 50% of population is killed	(μg/L)
LC50_exp_time	ObsTElapsed	exposure time in toxicity determination	(h)
K2	K2	elimination rate constant	(1/d)
EC50_growth	EC50Growth	external concentration of toxicant at which there is a 50% reduction in growth	(µg/L)
Growth_exp_time	ObsTElapsed	exposure time in toxicity determination	(h)
EC50_repro	EC50Repro	external concentration of toxicant at which there is a 50% reduction in reprod	(µg/L)
Repro_exp_time	ObsTElapsed	exposure time in toxicity determination	(h)
Ave_wet_wt	WetWt	mean wet weight of organism	(g)
Lipid_frac	LipidFrac	fraction of lipid in organism	(g lipid/g organ
Drift_Thresh	Drift Threshold	concentration at which drift is initiated	(µg/L)
TPlantToxRecord	Plant Toxicity Parameter	For each Chemical Simulated, the following parameters are required for each pl	
EC50_photo	EC50Photo	external concentration of toxicant at which there is 50% reduction in photosyn	(μg/L)
EC50_exp_time	ObsTElapsed	exposure time in toxicity determination	(h)
K2	K2	elimination rate constant	(1/d)
LC50	LC50	external concentration of toxicant at which 50% of population is killed	(µg/L)
LC50_exp_time	ObsTElapsed	exposure time in toxicity determination	(h)
Lipid_frac	LipidFrac	fraction of lipid in organism	(g lipid/g organ
TChemical	Chemical Parameters	For each Chemical to be simulated, the following parameters are required	
InitialCond	Initial Condition	Initial Condition of the state variable	μg/L
Loadings	Inflow Loadings	Daily loading as a result of the inflow of water (excluding modeled upstream r	μg/L

INTERNAL	TECH DOC	DESCRIPTION	UNITS
Alt_Loadings[Pointso urce]	Point Source Loadings	Daily loading from point sources	(g/d)
Alt_Loadings[Direct Preci	Direct Precipitation Loa	Daily loading from direct precipitation	(g/m2·d)
Alt_Loadings[NonPointsour	Non-Point Source Loading	Daily loading from non-point sources	(g/d)
Tox_Air	Gas-phase concentration		(g/m3)
TRemineralize	Nutrient Parameters	For each Nutrient to be simulated, O2 and CO2, the following parameters are re	
InitialCond	Initial Condition	Initial Condition of the state variable	mg/L
Loadings	Inflow Loadings	Daily loading as a result of the inflow of water (excluding modeled upstream r	mg/L
Alt_Loadings[Pointso urce]	Point Source Loadings	Daily loading from point sources	(g/d)
Alt_Loadings[Direct Preci	Direct Precipitation Loa	Daily loading from direct precipitation	(g/m2·d)
Alt_Loadings[NonPointsour	Non-Point Source Loading	Daily loading from non-point sources	(g/d)
TSedDetr	Sed. Detritus Parameters	For the Labile and Refractory Sed. Detritus compartments, the following parameters	
InitialCond	Initial Condition	Initial Condition of the state variable	(g/m2)
TToxicant.InitialCond	Toxicant Exposure	Initial Toxicant Exposure of the state variable, for each chemical simulated	μg/kg
TDetritus	Susp & Dissolved Detritu	For the Suspended and Dissolved Detritus compartments, the following parameter	
		1 ,	
InitialCond	Initial Condition	Initial Cond. of susp. & diss. detritus, as organic matter, organic carbon, or	mg/L
Percent_Part_IC	Percent Particulate Init	Percent of Initial Condition that is particulate as opposed to dissolved detri	percentage
Percent_Refr_IC	Percent Refractory Init	Percent of Initial Condition that is refractory as opposed to labile detritus	percentage

INTERNAL	TECH DOC	DESCRIPTION	UNITS
Loadings	Inflow Loadings	Daily loading as a result of the inflow of water (excluding modeled upstream r	mg/L
Percent_Part	Percent Particulate Infl	Daily parameter; % of loading that is particulate as opposed to dissolved detr	percentage
Percent_Refr	Percent Refractory Inflo	Daily parameter; % of loading that is refractory as opposed to labile detritus	percentage
Alt_Loadings[Pointso urce]	Point Source Loadings	Daily loading from point sources	(g/d)
Percent_Part_PS	Percent Particulate Poin	Daily parameter; % of loading that is particulate as opposed to dissolved detr	percentage
Percent_Refr_PS	Percent Refractory Point	Daily parameter; % of loading that is refractory as opposed to labile detritus	percentage
Alt_Loadings[NonPointsour	Non-Point Source Loading	Daily loading from non-point sources	(g/d)
Percent_Part_NPS	Percent Particulate NonP	Daily parameter; % of loading that is particulate as opposed to dissolved detr	percentage
Percent_Refr_NPS	Percent Refractory NonPo	Daily parameter; % of loading that is refractory as opposed to labile detritus	percentage
TToxicant.InitialCond	Toxicant Exposure	Initial Toxicant Exposure of the state variable	µg/kg
TToxicant.Loads	Tox Exposure of Inflow L	Daily parameter; Tox. Exposure of each type of inflowing detritus, for each ch	μg/kg
TBuried Detritus	Buried Detritus	For Each Layer of Buried Detritus, the following parameters are required	
InitialCond	Initial Condition	Initial Condition of the state variable	(g/m2)
TToxicant.InitialCond	Toxicant Exposure	Initial Toxicant Exposure of the state variable, for each chemical simulated	µg/kg
TPlant	Plant Parameters	For each plant type simulated, the following parameters are required	
InitialCond	Initial Condition	Initial Condition of the state variable	mg/L
Loadings	Inflow Loadings	Daily loading as a result of the inflow of water (excluding modeled upstream r	mg/L
TToxicant.InitialCond	Toxicant Exposure	Initial Toxicant Exposure of the state variable	µg/kg
TToxicant.Loads	Tox Exposure of Inflow L	Daily parameter; Tox. Exposure of the Inflow Loadings, for each chemical simul	µg/kg

INTERNAL	TECH DOC	DESCRIPTION	UNITS
TAnimal	Animal Parameters	For each animal type simulated, the following parameters are required	
InitialCond	Initial Condition	Initial Condition of the state variable	mg/L
Loadings	Inflow Loadings	Daily loading as a result of the inflow of water (excluding modeled upstream r	mg/L
TToxicant.InitialCond	Toxicant Exposure	Initial Toxicant Exposure of the state variable	μg/kg
TToxicant.Loads	Tox Exposure of Inflow L	Daily parameter; toxic exposure of the Inflow Loadings, for each chemical simulated	µg/kg
TrophIntArray.Pref	Prefprey, pred	for each prey-type ingested, a preference value within the matrix of preferences	(unitless)
TrophIntArray.ECoeff	EgestCoeff	for each prey-type ingested, the fraction of ingested prey that is egested	(unitless)
TVolume	Volume Parameters	For each segment simulated, the following water flow parameters are required	
InitialCond	Initial Condition	Initial Condition of the state variable	(m3)
InflowLoad	Inflow of Water	Inflow of water	(m3/d)
DischargeLoad	Discharge of Water	Discharge of water	(m3 /d)
Site Characteristics	Site Characteristics	The following characteristics are required	
Temperature	Temperature	Daily parameter; temperature of the segment; Optional, can use annual mean	(°C)
Wind	Wind	Daily parameter; wind velocity 10 m above the water; Optional, default time se	(m/s)
Light	Light	Daily parameter; avg. light intensity at segment top; Optional, can use annual	(ly/d)
Photoperiod	Photoperiod	Fraction of day with daylight; Optional, can be calculated from latitude	(hr/d)
рН	pH	Daily parameter; pH of the segment.	(pH)
Physical Geometry	Physical Geometry	For each segment simulated, the following physical geometry parameters are required	

INTERNAL	TECH DOC	DESCRIPTION	UNITS
Thickness	Segment Thickness	Thickness of the segment	m
Surface Area	Surface Area	Surface area of the segment	(m2)